=> D HIS

(FILE 'HOME' ENTERED AT 11:50:46 ON 04 DEC 2000)

	FILE '	HCAPL	S' ENTERED AT 11:50:54 ON 04 DEC 2000	
L1		13	LIQUID PHASE CARRIER	
L2		2	NUCLEIC ACID SOLUTION PHASE SYNTHESIS	
L3		1	L2 NOT L1	
L4		422	SOLUTION PHASE (3W) SYNTHESIS	
L5		1	SOLUTION PHASE BIOPOLYMER SYNTHESIS	
L6		0	L5 NOT L1	
L7		87	SOLUTION PHASE (4A) SYNTHESIS (4A) (BIOPOLYMER OR BIO POLYME	R
OR				
L8		77	(PREPAR? OR MANUF? OR PRODUC?) AND L7	
L9		87	SYNTHES? AND L7	
L10		426	(L1 OR L2 OR L4)(6A)(PREPAR? OR MANUF? OR PRODUC? OR	
SYNTH	ES?			
L11		79	L7 AND L10	

=> D BIB ABS 1-10

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ANSWER 1 OF 79 HCAPLUS COPYRIGHT 2000 ACS
ΑN
     2000:590846 HCAPLUS
DN
     133:310129
     Development of a Solution-Phase Synthesis of
TΙ
     Minor Groove Binding Bis-Intercalators Based on Triostin A Suitable for
     Combinatorial Synthesis
ΑU
     Boger, Dale L.; Lee, Jae Kyoo
     Department of Chemistry and The Skaggs Institute for Chemical Biology,
CS
     Scripps Research Institute, La Jolla, CA, 92037, USA
     J. Org. Chem. (2000), 65(19), 5996-6000
CODEN: JOCEAH; ISSN: 0022-3263
SO
PB
     American Chemical Society
DT
     Journal
LA
     English
AB
     The development of a soln. phase synthesis
     of a triostin A analog (azatriostin A) is disclosed which is suitable for
     the prepn. of combinatorial libraries enlisting only liq.-liq. acid/base
     extns. for the isolation and purifn. of all intermediates and the final
     product.
RE.CNT 38
RE
(2) Addess, K; Nucleic Acids Res 1994, V22, P5484 HCAPLUS
(3) Albericio, F; Synthesis 1987, P271 HCAPLUS
(4) Alfredson, T; Biopolymers 1991, V31, P1689 HCAPLUS
(5) Boger, D; Bioorg Med Chem 1998, V6, P1347 HCAPLUS
(6) Boger, D; Bioorg Med Chem Lett 1997, V7, P1903 HCAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 2 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     2000:557699 HCAPLUS
AN
DN
     133:267143
ΤI
     Solid-phase synthesis of chemotactic peptides using .alpha.-azido acids
ΑU
     Tornoe, Christian W.; Sengelov, Henrik; Meldal, Morten
CS
     Department of Chemistry, Carlsberg Laboratory, Copenhagen, DK-2500, Den.
     J. Pept. Sci. (2000), 6(7), 314-320
CODEN: JPSIEI; ISSN: 1075-2617
SO
PΒ
     John Wiley & Sons Ltd.
DT
     Journal
LA
     English
     Four chemotactic peptides, For-Met-Xxx-Phe-OMe (Xxx = Aib, Deg, Dpg, or
AB
     Dph, where Aib = 2-aminoisobutyric acid, Deg = diethylglycine, Dpg =
     dipropylglycine, Dpg = diphenylglycine) with an .alpha.,.alpha.-
     disubstituted amino acid at position 2 have been synthesized by the azido
     acid method on solid-phase, and were tested for biol. activity. Dpg in
     the central position was found to be as active as the natural chemotactic
     peptide for chemotactic activity toward human neutrophils. Higher yields
     were obtained than previously reported soln.-phase
     syntheses of chemotactic peptides, and EEDQ
     (2-ethoxy-1-ethoxycarbonyl-1,2-dihydroquinoline) was used successfully
     the difficult solid-phase formylation of amino groups.
RE.CNT 16
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(3) Blankemeyer-Menge, B; Tetrahedron Lett 1990, V31, P1701 HCAPLUS

(4) Carpino, L; J Am Chem Soc 1993, V115, P4397 HCAPLUS

(6) Dentino, A; J Biol Chem 1991, V266, P18460 HCAPLUS

(7) Kent, S; Ann Rev Biochem 1988, V57, P957 HCAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 3 OF 79 HCAPLUS COPYRIGHT 2000 ACS

AN 2000:514138 HCAPLUS

DN 133:252720

TI An unnatural amino acid that mimics a tripeptide .beta.-strand and forms .beta.-sheet-like hydrogen-bonded dimers

AU Nowick, James S.; Chung, De Michael; Maitra, Kalyani; Maitra, Santanu; Stigers, Kimberly D.; Sun, Ye

CS Department of Chemistry, University of California, Irvine, CA, 92697-2025,

USA

SO J. Am. Chem. Soc. (2000), 122(32), 7654-7661 CODEN: JACSAT; ISSN: 0002-7863

PB American Chemical Society

DT Journal

LA English

GΙ

AB Unnatural amino acid 5-HO2CCONH-2-MeO-C6H3-CONHNH2 (I; abbreviated Hao) contains hydrazine, 5-amino-2-methoxybenzoic acid and oxalic acid, and it duplicates the hydrogen-bonding functionality of one edge of a tripeptide .beta.-strand. The 2,7-di(tert-butyl)fluorenylmethyloxycarbonyl (Fmoc*)-and tert-butyloxycarbonyl (Boc)-protected derivs. of Hao are prepd. efficiently and in high yields by the condensation of suitably protected derivs. of hydrazine, 5-amino-2-methoxybenzoic acid and oxalic acid. Fmoc*-Hao and Boc-Hao behave like typical Fmoc- and Boc-protected amino acids and can be incorporated into peptides by std. solid- and soln.-phase peptide synthesis

Ι

ΙI

techniques using carbodiimide coupling agents. Hao-contg. peptide Me2CHCO-Phe-Hao-Val-NHBu (II) forms a .beta.-sheetlike hydrogen-bonded dimer in CDC13 and CD3OD-CDC13 solns. Peptides contg. Hao and natural amino acids display hydrogen-bonding surfaces that are complementary to the hydrogen-bonding edges of protein .beta.-sheets.

RE.CNT 54

RE

(1) Abbenante, G; J Am Chem Soc 1995, V117, P10220 HCAPLUS

(2) Albericio, F; Int J Pept Protein Res 1987, V30, P206 HCAPLUS Searched by John Dantzman 703-308-4488

- (3) Albericio, F; J Org Chem 1990, V55, P3730 HCAPLUS
- (4) Alsina, J; Chem Eur J 1999, V5, P2787 HCAPLUS
- (5) Beijer, F; Angew Chem, Int Ed 1998, V37, P75 HCAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 4 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 2000:405663 HCAPLUS
- DN 133:223039
- TI Total Synthesis of Distamycin A and 2640 Analogs: A Solution-Phase Combinatorial Approach to the Discovery of New, Bioactive DNA Binding Agents and Development of a Rapid, High-Throughput Screen for Determining Relative DNA Binding Affinity or DNA Binding Sequence Selectivity
- AU Boger, Dale L.; Fink, Brian E.; Hedrick, Michael P.
- CS Department of Chemistry and The Skaggs Institute for Chemical Biology,

The

- Scripps Research Institute, La Jolla, CA, 92037, USA
- SO J. Am. Chem. Soc. (2000), 122(27), 6382-6394
- CODEN: JACSAT; ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 133:223039

GI

AB The development of a soln.-phase synthesis

Searched by John Dantzman 703-308-4488

of distamycin A and its extension to the prepn. of 2640 analogs are described. Thus, soln.-phase synthesis

techniques with reaction workup and purifn. employing acid/base lig.-liq. extns. were used in the multistep prepn. of distamycin A (8 steps, 40% overall yield) and a prototypical library of 2640 analogs providing intermediates and final products that are .gtoreq.95% pure on conventional

reaction scales. The complementary development of a simple, rapid, and high-throughput screen for DNA binding affinity based on the loss of fluorescence derived from displacement of prebound ethidium bromide is disclosed which is applicable for assessing relative or abs. binding affinity to DNA homopolymers or specific sequences (hairpin oligonucleotides). Using hairpin oligonucleotides, this method permits the screening of a library of compds. against a single predefined sequence

to identify high affinity binders, or the screening of a single compd. against a full library of individual hairpin oligonucleotides to define its sequence selectivity. The combination permits the establishment of the complete DNA binding profile of each member of a library of compds. Screening the prototypical library provided compds. that are 1000 times more potent than distamycin A in cytotoxic assays (I, Boc = tert-butoxycarbonyl; IC50 = 29 nM, L1210), that bind to poly[dA]-poly[dT] with comparable affinity, and that exhibit an altered DNA binding sequence

selectivity. Several candidates were identified which bound the five-base-pair AT-rich site of the PSA-ARE-3 sequence, and one (II, R =4-dimethylaminobutyryl; K = 3.2 .times. 106 M-1) maintained the high affinity binding (K = 4.5 .times. 106 M-1) to the ARE-consensus sequence contg. a GC base-pair interrupted five-base-pair AT-rich site suitable

inhibition of gene transcription initiated by hormone insensitive androgen

receptor dimerization and DNA binding characteristic of therapeutic resistant prostate cancer.

RE.CNT 55 RE

for

- (1) Abu-Daya, A; Nucleic Acids Res 1995, V23, P3385 HCAPLUS
- (2) Abu-Daya, A; Nucleic Acids Res 1997, V25, P4962 HCAPLUS
- (4) Baguley, B; Nucleic Acids Res 1978, V5, P161 HCAPLUS(5) Baird, E; J Am Chem Soc 1996, V118, P6141 HCAPLUS
- (6) Behrens, C; Comb Chem High Throughput Screening 1998, V1, P127 HCAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11ANSWER 5 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- 2000:331625 HCAPLUS AN
- TΙ Identification of new chemical motifs that bind the a-site subdomain of 16S ribosomal RNA using solution-phase combinatorial library synthesis techniques.
- Kung, Pei-Pei; Lowery, Kristin; Wheeler, Patrick; Hofstadler, Steven; ΑU Swayze, Eric; Griffey, Richard
- CS
- Ibis Therapeutics, Carlsbad, CA, 92008, USA Book of Abstracts, 219th ACS National Meeting, San Francisco, CA, March SO 26-30, 2000 (2000), MEDI-025 Publisher: American Chemical Society, Washington, D. C. CODEN: 69CLAC
- Conference; Meeting Abstract DT
- LA English

AB The use of soln. phase combinatorial library synthesis techniques and simultaneous addn. of functionalities enabled us to efficiently prep. combinatorial libraries with diverse structures which possess potential RNA-binding motifs. The technique of simultaneous addn. of stoichiometric amts. of coupling reagents was used to attach functionalities to several sym. or asym. bi-functional scaffolds

utilizing alkylation, acylation, and amidation reactions. Support-bound bases, catalysts, as well as scavengers were used to perform the alkylation reactions, the acylation reactions with isocyanates, and the HATU-activated amidation reactions. The chem. identities and 16S RNA binding activities of the combinatorial mols. were detd. by mass spectrometry.

- L11 ANSWER 6 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 2000:288369 HCAPLUS
- DN 133:53934
- TI Synthetic agouti protein fragment (91-131) is an inverse agonist of the melanocortin-1 (MC-1) receptor
- Eberle, Alex N.; Froidevaux, Sylvie; Meier, Maja; Jaggin, Verena; Bodi, ΑU Jozsef; Orosz, Gyorgy; Suli-Vargha, Helga
- Department of Research (ZLF), University Hospital and University CS
- Children's Hospital, Basel, CH-4031, Switz.
 Pept. 1998, Proc. Eur. Pept. Symp., 25th (1999), Meeting Date 1998, SO 66-67.

Editor(s): Bajusz, Sandor; Hudecz, Ferenc. Publisher: Akademiai Kiado, Budapest, Hung. CODEN: 68WKAY

- DT Conference
- LA English
- AB To obtain more information about the biol. characteristics of the C-terminal part of the agouti protein (AP) at the melanocortin-1 (MC-1) receptor, the authors studied the synthetic (91-131) AP fragment using mouse and human melanoma cells. The chem. synthesis of the C-terminal (91-131) region of AP was performed by combination of solid phase and soln. phase peptide synthesis. The

biol. characterization of AP(91-131) was carried out with four different assay systems, namely, the MC-1 receptor binding assay, the adenylate cyclase assay, the tyrosinase assay and the melanin assay. In the binding

assay, the potency of the AP(91-131) fragment as a competitor of .alpha.-MSH was only 56% compared to that of AP. In the tyrosinase and melanin assays, AP(91-131) was also less potent than AP(1-131). The agouti fragment, however, inhibited basal adenylate cyclase activity in B16-F1 cell membranes more effectively than the intact agouti protein.

In summary, AP(91-131) displays the same biol. characteristics found with AP:

it antagonizes .alpha.-MSH binding to MC-1 receptors and signaling in B16-F1 cells at the level of adenylate cyclase, tyrosinase and melanogenesis. However, the fact that AP(91-131) reduces basal cellular cyclase, tyrosinase and melanogenic activity in unstimulated B16-F1

indicates that AP(91-131) is an inverse agonist with similar characteristics and even higher potency (in the adenylate cyclase assay) than the parent full-length agouti protein.

RE.CNT

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RE
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(2) Bodi, J; Tetrahedron Lett 1997, V38, P3293 HCAPLUS
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(4) Chhajlani, V; FEBS Lett 1992, V309, P417 HCAPLUS
(7) Lu, D; Nature 1994, V371, P799 HCAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 7 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     2000:288367 HCAPLUS
ΑN
DN
     133:105311
TΙ
     Statistical combination of thymus peptides, a synthetic library mimicking
     the physiological environment
     Birr, Christian; Braum, Gunther; Hirt, Werner; Klett-Loch, Gunther H.
ΑU
     Faculty of Chemistry, Heidelberg University, Heidelberg, D-69120, Germany
CS
     Pept. 1998, Proc. Eur. Pept. Symp., 25th (1999), Meeting Date 1998,
SO
62-63.
     Editor(s): Bajusz, Sandor; Hudecz, Ferenc. Publisher: Akademiai Kiado,
     Budapest, Hung.
     CODEN: 68WKAY
DT
     Conference
LA
     English
AB
     A symposium report. We have synthesized a statistical chem. library of
     thymus peptides by employing stepwise soln.
     phase peptide synthesis conditions on those
     amino acids characteristic in quantity and nature to thymus tissue
     hydrolyzates.
    ANSWER 8 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     2000:234109 HCAPLUS
AN
DN
     132:334780
ΤI
     Solution synthesis of peptides
ΑU
     Sakakibara, Shumpei
     Protein Research Foundation, Peptide Institute, Inc., Osaka, 562, Japan
CS
     Collect. Symp. Ser. (1999), 1(Future Aspects in Peptide Chemistry), 1-11
SO
     CODEN: CSYSFN
PB
     Institute of Organic Chemistry and Biochemistry, Academy of Sciences of
     the Czech Republic
DT
     Journal
     English
LA
     A symposium on the author's work, comparing the effectiveness of
AB
     soln. phase synthesis of peptides to
     solid-phase peptide synthesis.
RE.CNT 37
(4) Chino, N; Biochem Biophys Res Commun 1988, V151, P1285 HCAPLUS
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(9) Kimura, T; Biochem Biophys Res Commun 1983, V114, P493 HCAPLUS
(10) Kimura, T; Biochem Soc Trans 1990, V18, P1297 HCAPLUS
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ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 9 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
AN
     2000:176120 HCAPLUS
DN
     133:4948
     Solution-Phase Synthesis of a Hindered
ΤI
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N-Methylated Tetrapeptide Using Bts-Protected Amino Acid Chlorides: Searched by John Dantzman 703-308-4488

Efficient Coupling and Methylation Steps Allow Purification by Extraction ΑU Vedejs, Edwin; Kongkittingam, Chutima CS Department of Chemistry, University of Michigan, Ann Arbor, MI, 48109, USA J. Org. Chem. (2000), 65(8), 2309-2318 CODEN: JOCEAH; ISSN: 0022-3263 SO PB American Chemical Society DT Journal LA English AB N-Benzothiazole-2-sulfonyl (Bts)-protected amino acid chlorides were used to prep. the hindered cyclosporin 8-11 tetrapeptide subunit. synthesis was performed via 3a and the deprotected amines (S)-MeVal-OCMe3, (S)-MeLeu-(S)-MeVal-OCMe3, and (S)-MeLeu-(S)-MeLeu-(S)-MeVal-OCMe3, including three repeated cycles involving N-methylation with MeI-K2CO3, deprotection of the Bts group, and N-acylation with an N-Bts-amino acid chloride. Among three Bts cleavage methods compared (H3PO2-THF, NaBH4-EtOH, PhSH-K2CO3), the third gave somewhat higher overall yields. N-Acylation of (S)-MeVal-OCMe3 with Bts-protected N-methylamino acid chloride followed by deprotection was also highly efficient and could be used as an alternative route to Bts-(S)-MeLeu-(S)-MeVal-OCMe3. Each of the deprotected amines was isolated without chromatog. using simple extn. methods to remove neutral byproducts. The tetrapeptide was obtained in anal. pure form as the monohydrate. RE.CNT 21 RE (1) Akaji, K; J Org Chem 1999, V64, P405 HCAPLUS (2) Boger, D; J Am Chem Soc 1998, V120, P7220 HCAPLUS (3) Bowman, W; Tetrahedron 1997, V53, P15787 HCAPLUS (4) Carpino, L; Acc Chem Res 1996, V29, P268 HCAPLUS (5) Carpino, L; Tetrahedron Lett 1998, V39, P241 HCAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT L11 ANSWER 10 OF 79 HCAPLUS COPYRIGHT 2000 ACS 1999:775928 HCAPLUS AN DN 132:103146 TI Stimulation of nonspecific resistance by thymopentin and its analogs against Leishmania donovani infection in hamsters Sharma Anuradha, P.; Rohatgi, A.; Haq, W.; Mathur, K. B.; Katiyar, J. C. ΑU Divisions of Parasitology and Biopolymers, Central Drug Research CS Institute, Lucknow, India Peptides (N. Y.) (1999), 20(11), 1381-1383 SO CODEN: PPTDD5; ISSN: 0196-9781 PB Elsevier Science Inc. DTJournal LA English AB Thymopentin and its analogs have been synthesized by the soln. phase method of peptide synthesis and evaluated for their prophylactic efficacy against L. donovani infection in hamsters. Thymopentin and some of the analogs were found to stimulate nonspecific resistance of the host against leishmania donovani infection in hamsters. RE.CNT RE

- (1) Audhya, T; Proc Natl Acad Sci 1984, V81, P2847 HCAPLUS
- (2) Cordero, O; Immunol Today 1997, V18, P10 HCAPLUS
- (3) Diezel, W; Int J Immunopharmacol 1993, V15, P269 HCAPLUS Searched by John Dantzman 703-308-4488

- (5) Goldstein, A; Biological response modifiers in the treatment of cancer and infectious diseases 1993, P39 HCAPLUS
- (7) Rastogi, A; FEBS Lett 1993, V317, P93 HCAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> D BIB ABS 11-79

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ANSWER 11 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
AN
     1999:708779 HCAPLUS
DN
     131:351620
TΙ
     Solution phase biopolymer synthesis
     of oligodeoxyribonucleotides using multifunctional lig
      . phase carriers
     Koster, Hubert; Worl, Ralf
IN
PA
     USA
SO
     PCT Int. Appl., 88 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
                         KIND
                                DATE
     PATENT NO.
                                                 APPLICATION NO.
                                                                     DATE
                         ____
                                -----
PΙ
     WO 9955718
                          A2
                                19991104
                                                 WO 1999-US8939
                                                                     19990426
     WO 9955718
                          Α3
                                19991216
          W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
              DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU,
               TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     AU 9936643
                          A1 19991116
                                                 AU 1999-36643
                                                                     19990426
PRAI US 1998-67337
                         19980427
     WO 1999-US8939
                         19990426
     Multifunctional liq. phase carriers (LPCs) and methods of using LPCs for
AB
     the prepn. of biopolymers are provided. The LPCs are highly sym. compds.
     that possess more than two points of attachment for biopolymer synthesis.
     The LPCs have the formula Sp(X1)n, where Sp is a highly sym. moiety such
     that all X1 groups are equiv. X1 is a functional group that is suitable
     for biopolymer synthesis, including OH, SH, NH2, COOH and the like.
     Biopolymers that may be produced using the methods provided include
     oligonucleotides, peptides, protein nucleic acids (PNAs) and
     oligosaccharides. Analogs of the biopolymers may also be prepd. using
the
     methods. Thus decamer d(GACCGGCAGT) was prepd. using
     multifunctional liq. phase carriers.
     ANSWER 12 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     1999:442438 HCAPLUS
AN
DN
     131:239827
     Radiometal-labelled macrocyclic chelator-derivatized somatostatin
TΙ
analogue
     with superb tumour-targeting properties and potential for
     receptor-mediated internal radiotherapy
     Heppeler, A.; Froidevaux, S.; Macke, H. R.; Jermann, E.; Behe, M.;
Powell,
     P.; Hennig, M.
CS
     Institute of Nuclear Medicine, Div. of Radiological Chemistry, University
                      Searched by John Dantzman 703-308-4488
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Hospital Basel, Basel, CH-4031, Switz.
    Chem.--Eur. J. (1999), 5(7), 1974-1981
CODEN: CEUJED; ISSN: 0947-6539
SO
PΒ
    Wiley-VCH Verlag GmbH
DT
     Journal
LA
     English
AB
    A monoreactive DOTA (1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic
     acid) prochelator (4,7,10-tricarboxymethyl-tert-Bu ester
     1,4,7,10-tetraazacyclododecane-1-acetate) was synthesized which is useful
     in solid-phase and soln.-phase peptide
     synthesis; it was coupled to the somatostatin analog
     Tyr3-Lys5(BOC)-octreotide. Deprotection in one step afforded
     DOTA0-D-Phel-Tyr3-octreotide (DOTATOC) in .apprxeq.65% yield.
     peptide, modified with a chelator, was complexed with the radiometals
     67Ga3+, 111In3+ and 90Y3+ in high yields and with high specific
     activities. The three radiopeptides show high stability in human serum
     and high affinity to the somatostatin receptor: it is four to five times
     higher for 67Ga-DOTATOC compared to 90Y-DOTATOC and 111In-DOTATOC. The
     67Ga-labeled compd. also shows significantly higher tumor and lower
kidnev
     uptake than the two congeners. 67Ga-DOTATOC was compared in patients
with
     the com. available gold std. 111In-DTPA0-D-Phel-octreotide.
     compd. delineates SRIF-receptor pos. tumors very favorably and shows
     distinctly lower uptake by the kidneys. Evidently, the differences in
the
     coordination chem. of the metals causes the differences in the biol.
     behavior. Indeed, a crystallog. study of the Ga3+ and Y3+ complexes of
     the model peptide DOTA-D-PheNH2 showed differences in the geometry of the
     complexes. The gallium complex is hexacoordinated with pseudooctahedral
     cis geometry and a folded macrocyclic unit. The equatorial plane is
     formed by two transannular nitrogens of the cyclen ring and two oxygens
of
     the corresponding carboxylate groups. The two axial positions are formed
    by the two remaining ring nitrogen atoms. The amide carboxy oxygen is
not
     bound to the metal and one carboxylate group is free and most likely
     contributes to the favorable handling of the radiopeptide by the kidneys.
     In contrast, the structure of Y-DOTA-D-PheNH2 has eight-fold
coordination,
     and includes the amide carboxy oxygen. The geometry is a compact and
     somewhat distorted square-antiprism with two almost perfect planes (N4
and
    O4) with a max. deviation of 0.025 A. The dihedral angle between the two
    planes is only 0.36.degree..
RE.CNT 48
(2) Aime, S; Angew Chem Int Ed 1998, V37, P2673 HCAPLUS
(3) Aime, S; Chem Soc Rev 1998, V27, P19 HCAPLUS
(4) Aime, S; Inorg Chem 1992, V31, P4291 HCAPLUS
(5) Albert, R; Actualite de Chimie Therapeutique 1994, V21, P111 HCAPLUS
(6) Albert, R; Bioorg Med Chem Letters 1998, V8, P1207 HCAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
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Searched by John Dantzman 703-308-4488

L11 ANSWER 13 OF 79 HCAPLUS COPYRIGHT 2000 ACS

1999:188594 HCAPLUS

131:19271

AN DN

```
ΤI
     Convergent solution-phase synthesis of a
     nucleopeptide using a protected oligonucleotide
ΑU
     McMinn, Dustin L.; Greenberg, Marc M.
CS
     Department of Chemistry, Colorado State University, Fort Collins, CO,
     80523, USA
SO
     Bioorg. Med. Chem. Lett. (1999), 9(4), 547-550
     CODEN: BMCLE8; ISSN: 0960-894X
PB
     Elsevier Science Ltd.
DT
     Journal
LA
     English
AB
     A nucleopeptide was prepd. in a convergent manner via segmental coupling
     of the protected biopolymers in soln. The resulting nucleopeptide contg.
     the binding site of .lambda. repressor and a peptide contg. the consensus
     sequence of the DNA binding helix of the helix turn-helix-proteins was
     obtained in 72% yield using only five equiv. of the peptide relative to
     the oligonucleotide. This demonstrates that the recently developed
method
     for the soln. phase coupling of protected oligonucleotides is amenable to
     the convergent synthesis of larger nucleopeptides that are potentially
     capable of adopting secondary structure.
RE.CNT 20
RE
(1) Bergmann, F; Tetrahedron Lett 1995, V36, P1839 HCAPLUS
(3) de la Torre, B; Tetrahedron Lett 1994, V35, P2733 HCAPLUS
(4) Erout, M; Bioconjugate Chem 1996, V7, P568 HCAPLUS(5) Jones, D; Bioconjugate Chem 1994, V5, P390 HCAPLUS
(6) Kahl, J; J Org Chem 1998, V63, P4870 HCAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 14 OF 79 HCAPLUS COPYRIGHT 2000 ACS
AN
     1999:139856 HCAPLUS
DN
     130:153924
TΙ
     Solution phase synthesis of oligonucleotides
IN
     Reese, Colin Bernard; Song, Quanlai
PA
     Zeneca Limited, UK
SO
     PCT Int. Appl., 32 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
     PATENT NO.
                       KIND DATE
                                            APPLICATION NO.
                                                              DATE
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ΡI
     WO 9909041
                       A2
                             19990225
                                             WO 1998-GB2407
                                                               19980810
     WO 9909041
                       А3
                             19990506
         W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
             DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG,
             KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX,
             NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT,
             UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI,
             CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                            AU 1998-87386
     AU 9887386
                       A1
                             19990308
                                                               19980810
     EP 1003758
                       A2
                             20000531
                                           EP 1998-938782
                                                               19980810
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
     NO 2000000690
                                            NO 2000-690
                            20000411
                                                               20000211
                       Α
                    Searched by John Dantzman 703-308-4488
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PRAI GB 1997-17158
                       19970813
     WO 1998-GB2407
                       19980810
     MARPAT 130:153924
OS
     A process for the synthesis in soln. phase of a phosphorothicate triester
AB
     is provided. The process comprises the soln. phase coupling of an
     H-phosphonate with an alc. in the presence of a coupling agent to form an
     H-phosphonate diester. The H-phosphonate diester is oxidized in situ
with
     a sulfur transfer agent to produce the phosphorothioate triester.
     Preferably, the H-phosphonate and alc. are protected nucleosides or
     oligonucleotides. Oligonucleotide H-phosphonates which can be used in
the
     formation of phosphorothioate triesters are also provided.
     ANSWER 15 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     1999:98326 HCAPLUS
AN
DN
     130:196945
ΤI
     Solution phase synthesis of potential
     DNA-binding molecules based on the PNA backbone
ΑU
     Challa, Hemavathi; Woski, Stephen A.
CS
     Department of Chemistry and Coalition for Biomolecular Products, The
     University of Alabama, Tuscaloosa, AL, 35487-0336, USA
SO
     Tetrahedron Lett. (1999), 40(3), 419-422
     CODEN: TELEAY; ISSN: 0040-4039
PB
     Elsevier Science Ltd.
DT
     Journal
     English
LA
AΒ
     The N-(2-aminoethyl)glycine backbone unit of PNA has been derivatized
with
     pyrene-acetic acid and acetic acid moieties to produce monomers for the
     synthesis of potential poly-intercalators. Short oligomers contg. these
     residues have been assembled using soln. phase coupling reactions.
RE.CNT 22
RE
(1) Armitage, B; Nucleic Acids Res 1998, V26, P715 HCAPLUS(2) Armitage, B; Proc Natl Acad Sci USA 1997, V94, P12320 HCAPLUS
(3) Atwell, G; J Med Chem 1986, V29, P69 HCAPLUS
(4) Chen, F; Nucleic Acids Res 1983, V11, P7231 HCAPLUS
(6) Dueholm, K; New J Chem 1997, V21, P19 HCAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 16 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
ΑN
     1998:804800 HCAPLUS
DN
     130:153914
ΤI
     Solution-Phase Bioconjugate Synthesis Using
     Protected Oligonucleotides Containing 3'-Alkyl Carboxylic Acids
AU
     Kahl, Jeffrey D.; Greenberg, Marc M.
     Department of Chemistry, Colorado State University, Fort Collins, CO,
CS
     80523, USA
SO
     J. Org. Chem. (1999), 64(2), 507-510
     CODEN: JOCEAH; ISSN: 0022-3263
PB
     American Chemical Society
DT
     Journal
LA
     English
     Protected oligonucleotides contg. 3'-alkyl carboxylic acids are obtained
AB
     from a photolabile solid-phase synthesis support (1b). The protected
     oligonucleotides are efficiently conjugated (>80%) with amines in soln.
to
                    Searched by John Dantzman 703-308-4488
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yield products of high purity under mild reaction conditions. This method

is particularly well-suited for the synthesis of oligonucleotide-peptide conjugates contg. a covalent linkage between the 3' terminus of an oligonucleotide and the amino terminus of a peptide. High yields of nucleopeptides are obtained even when the peptide contains a hindered N-terminal amino acid.

RE.CNT 24

RE

- (1) Beaucage, S; Tetrahedron 1993, V49, P1925 HCAPLUS(2) Beaucage, S; Tetrahedron 1993, V49, P6123 HCAPLUS
- (3) Bischoff, R; Anal Biochem 1987, V164, P336 HCAPLUS
- (4) Erout, M; Bioconjugate Chem 1996, V7, P568 HCAPLUS
- (5) Ghosh, S; Nucleic Acids Res 1987, V15, P5353 HCAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT
- ANSWER 17 OF 79 HCAPLUS COPYRIGHT 2000 ACS L11
- AN 1998:667156 HCAPLUS
- DN 130:4017
- TΙ A new approach to oligonucleotide synthesis in solution
- ΑU Reese, Colin B.; Song, Quanlai
- CS Department of Chemistry, King's College London, London, WC2R 2LS, UK
- SO Nucleosides Nucleotides (1998), 17(9-11), 2027-2031 CODEN: NUNUD5; ISSN: 0732-8311
- Marcel Dekker, Inc. PB
- Journal DT
- LA English
- AΒ A symposium on new approach, based on the use of 3'-H-phosphonate

blocks, is described for the synthesis of oligodeoxyribonucleotides and their phosphorothioate analogs in soln.

RE.CNT 16

- (1) Beaucage, S; Methods in Molecular Biology Vol 20 Protocols for Oligonucleotides and Analogs 1993, P33 HCAPLUS
- (2) Behforouz, M; J Org Chem 1969, V34, P51 HCAPLUS
- (3) Chattopadhyaya, J; Nucleic Acids Res 1980, V8, P2039 HCAPLUS
- (4) Froehler, B; Methods in Molecular Biology Vol 20 Protocols for Oligonucleotides and Analogs 1993, P63 HCAPLUS
- (5) Gura, T; Science 1995, V270, P575 HCAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 18 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1998:645441 HCAPLUS
- DN 130:25282
- ΤI The asymmetric synthesis of arginine mimetics: derivatives of (S)-2-, 3and 4-amidinophenylalanine suitable for incorporation into enzyme inhibitors and/or peptides
- Kent, D. R.; Cody, W. L.; Doherty, A. M. ΑU
- Parke-Davis Pharmaceutical Research, Division of Warner-Lambert Company, CS Ann Arbor, MI, USA
- J. Pept. Res. (1998), 52(3), 201-207 SO CODEN: JPERFA; ISSN: 1397-002X
- PΒ Munksgaard International Publishers Ltd.
- DT Journal
- LA English
- Ortho, meta and para isomers of amidinophenylalanine represent modified AB Searched by John Dantzman 703-308-4488

arginine residues and are important synthetic intermediates for enzyme inhibitors. Thus, a convenient asym. synthesis of (S)-N.alpha.-(tert-butyloxycarbonyl)-2-, (S)-N.alpha.-(tert-butyloxycarbonyl)-3-, and (S)-N.alpha.-(tert-butyloxycarbonyl)-4-amidinophenylalanine N,O-dimethylamides (Weinreb amides) is described here. These derivs. represent key synthetic intermediates for the synthesis of enzyme inhibitors because the amidino moiety can be readily orthogonally protected, while the Weinreb amide is easily converted to a variety of electrophilic carbonyls via redn. to the corresponding aldehyde or by reaction with various lithiated heterocycles. Also, the Weinreb amide

can

be reduced to the aldehyde and subsequently oxidized to the corresponding carboxylate, which is suitable for solid- or soln.-phase peptide synthesis.

RE.CNT 15

RE

- (1) Bergner, A; J Enzyme Inhib 1995, V9, P101 HCAPLUS
- (2) Das, J; Bioorg Med Chem 1995, V3, P999 HCAPLUS
- (3) Dickneite, G; Thromb Res 1995, V77, P357 HCAPLUS
- (4) Edmunds, J; Annual Reports in Medicinal Chemistry 1996, P51 HCAPLUS
- (5) Fehrentz, J; Synthesis 1983, P676 HCAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 19 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1998:500424 HCAPLUS
- DN 129:260748
- TI Chemical synthesis of peptides
- AU Hruby, Victor J.; Meyer, Jean-Philippe
- CS University of Arizona, USA
- SO Bioorg. Chem.: Pept. Proteins (1998), 27-64, 473-479. Editor(s): Hecht, Sidney M. Publisher: Oxford University Press, New York, N. Y. CODEN: 66LQAH
- DT Conference; General Review
- LA English
- AB A review with 242 refs. providing an overview of the synthetic methodol. available both for soln. phase peptide synthesis and solid phase peptide synthesis.

The review emphasizes general considerations that are important in peptide

synthesis, introduces current topics of general interest, and points to more comprehensive treatments and other aspects of the subject in the literature.

- L11 ANSWER 20 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1998:490653 HCAPLUS
- DN 129:136440
- TI Product anchored sequential synthesis method for solution phase prepn. of oligonucleotides and peptides
- IN Pieken, Wolfgang; Gold, Larry
- PA Nexstar Pharmaceuticals, Inc., USA
- SO PCT Int. Appl., 104 pp. CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 2

PATENT NO. KIND DATE APPLICATION NO. DATE Searched by John Dantzman 703-308-4488

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ΡI
      WO 9830578
                           A1
                                 19980716
                                                   WO 1998-US562
                                                                        19980106
          W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
               DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG,
               KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX,
          NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI,
               FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM,
               GA, GN, ML, MR, NE, SN, TD, TG
      US 5874532
                                 19990223
                                                   US 1997-780517
                                                                        19970108
                           Α
      AU 9860223
                           Α1
                                 19980803
                                                   AU 1998-60223
                                                                        19980106
      EP 996627
                           Α1
                                 20000503
                                                   EP 1998-903457
                                                                        19980106
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
               IE, FI
PRAI US 1997-780517
                          19970108
      WO 1998-US562
                          19980106
os
      MARPAT 129:136440
GI
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AB This invention discloses an improved method (product anchored sequential synthesis, PASS) for the sequential soln. phase synthesis of oligonucleotides and peptides via selective retention of protected lipophilic intermediates on a C18 resin, or by covalent (Diels-Alder reaction) attachment of diene-contg. protected intermediates to dienophile-derivatized resins. The method lends itself to automation and is ideally suited for large scale manuf. oligonucleotides with high efficiency. Thus, diene-contg. protected monomer I, prepd. in several steps from 3,5-hexadien-1-ol, 4,4'-dihydroxyacetophenone, PhMgBr, thymidine, and (Me2CH)2NP(Cl)OCH2CH2CN, was coupled with a polyethylene glycol (PEG) thymidine deriv., anchored to a maleimide-derivatized polystyrene resin Searched by John Dantzman 703-308-4488

via a Diels-Alder reaction, purified, and cleaved to yield pure polyethylene glycol-derivatized dimer PEG-dTdT-OH.

- L11 ANSWER 21 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1998:395314 HCAPLUS
- DN 129:161833
- TI Use of 1-.beta.-naphthalenesulfonyloxybenzotriazole as coupling reagent for peptide synthesis in the presence of fluorinated alcohols as cosolvent
- AU Khare, Sanjay K.; Singh, Geeta; Agarwal, Kamlesh C.; Kundu, Bijoy CS Division of Bioploymers, Central Drug Research Institute, Lucknow, 226001.

India

- SO Protein Pept. Lett. (1998), 5(3), 171-174
 - CODEN: PPELEN; ISSN: 0929-8665
- PB Bentham Science Publishers
- DT Journal
- LA English
- AB Soln. phase synthesis of peptides

in solvents mixed with fluorinated alcs. have been carried out using 1-.beta.-naphthalenesulfonyloxybenzotriazole (NSBt) as coupling reagent.

- L11 ANSWER 22 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1998:391150 HCAPLUS
- DN 129:149224
- TI Reactivity and suitability of t-Boc-protected thiophosphotyrosine intermediate analogs for the solid or solution phase peptide synthesis
- AU Kim, Eun-Kyung; Choi, Heesung; Lee, Eung-Seok
- CS College of Pharmacy, Yeungnam University, Kyongsan, 712-749, S. Korea
- SO Arch. Pharmacal Res. (1998), 21(3), 330-337 CODEN: APHRDQ; ISSN: 0253-6269
- PB Pharmaceutical Society of Korea
- DT Journal
- LA English
- OS CASREACT 129:149224

GΙ

AB Protected O-thiophosphono-L-tyrosine derivs. I (R = Me, CH2CH2CN; Boc = Me3CO2C) were prepd. as intermediates for the synthesis of thiophosphotyrosine-contg. peptides. The reactivity and suitability of two compds. for the solid phase or soln. phase peptide synthesis utilizing Boc chem. were examd.

- L11 ANSWER 23 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1998:159572 HCAPLUS
- DN 128:230678
- TI Application of AlMe3-mediated amidation reactions to solution phase peptide synthesis
- AU Martin, Stephen F.; Dwyer, Michael P.; Lynch, Christopher L.
- CS Department of Chemistry and Biochemistry, The University of Texas at Austin, Austin, TX, 78712, USA
- Austin, Austin, TX, 78712, USA SO Tetrahedron Lett. (1998), 39(12), 1517-1520 CODEN: TELEAY; ISSN: 0040-4039
- PB Elsevier Science Ltd.
- DT Journal
- LA English
- OS CASREACT 128:230678
- AB A practical modification of the Weinreb amidation protocol employing amino

acids as the amine reaction partner has been developed that allows for the

facile synthesis of oligopeptides in soln. Thus, treatment of an amino acid (or a dipeptide) with AlMe3 in 1,2-dichloroethane/hexane for 30 min, followed by addn. of an N-protected amino acid ester or an N-protected peptide ester gave the corresponding N-protected peptide in 31-60%

Similar reactions of amino acids with carboxylic acid esters or .beta.-butyrolactone gave N-acylated amino acid derivs. in 59-77% yields.

- L11 ANSWER 24 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1998:139768 HCAPLUS
- TI TFFH, a versatile reagent for organic transformations in solid- and solution-phase.
- AU Pillai, Sasi K.; Kates, Steven A.; Purkayastha, Subhasish
- CS PerSeptive Biosystems, Framingham, MA, 01701, USA
- SO Book of Abstracts, 215th ACS National Meeting, Dallas, March 29-April 2 (1998), ORGN-251 Publisher: American Chemical Society, Washington, D. C. CODEN: 65QTAA
- DT Conference; Meeting Abstract
- LA English
- AB Tetramethylfluoroformamidinium hexafluorophosphate (TFFH) is an effective activator recently introduced for both solid- and soln.-

phase peptide synthesis. TFFH converts

carboxylic acids to their corresponding acid fluorides, which are useful precursors for a variety of synthetic transformations. To explore the utility of this reaction in org. synthesis, apart from peptide assembly, several methods both in soln.— and solid—phase were examd. Thus, a

simple

and convenient one-pot conversion of carboxylic acids to alcs. was developed. A wide variety of acid substrates, including Fmoc- and Boc-protected amino acids, were reduced to the resp. alcs. in high yields and with retention of optical configuration. The protocol was also extended to the solid-phase construction of peptide-alcs. Similarly, one-pot procedures for the conversion of carboxylic acids to aldehydes, esters, amides, and thioesters; and sulfonic acids to sulfonamides, also were elaborated.

- L11 ANSWER 25 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1998:1337 HCAPLUS
- DN 128:75677

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ΤI
     Use of propylene oxide as an acid scavenger in peptide synthesis
     Dhaon, Madhup K.
IN
PA
     Abbott Laboratories, USA
     U.S., 4 pp.
SO
     CODEN: USXXAM
DΤ
     Patent
LA
     English
FAN.CNT 1
                  KIND DATE
                                          APPLICATION NO. DATE
     PATENT NO.
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                                                            -----
     US 5698676 A 19971216 US 1995-565465 19951130
PΙ
AB
     A process of using an alkylene oxide as an acid scavenger during
     peptide syntheses in both solid and soln.
     phases is claimed. The steps of this process include reacting an
     N.alpha.-Boc-protected amino acid with an N.alpha.-unprotected amino acid
     to form a peptide contg. Boc-protected amino terminus, deprotecting the
     formed peptide of the Boc group with an acid, and neutralizing the acid
     with an alkylene oxide soln. For example, to a soln. of Cbz-Phe-OBt (OBt = hydroxybenzotriazole ester) in THF/CH2Cl2 were added, in the given
     order, EDAC [1-ethyl-3-(3-dimethylaminopropyl)cabodiimide hydrochloride],
     H-Gly-OCMe3.cntdot.HCl, and a soln. of propylene oxide in THF. After a
     workup that included the addn. of HCl, the dipeptide, Cbz-Phe-Gly-OCMe3,
     was collected at an yield of 95%.
     ANSWER 26 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     1997:758228 HCAPLUS
AN
DN
     128:48457
ΤI
     Solution phase synthesis of an
     oligodeoxyribonucleotide phosphorothioate for therapeutic
     applications
     Cheruvallath, Z. S.; Krotz, A. H.; Cole, D. L.; Ravikumar, V. T.
ΑU
CS
     Isis Pharmaceuticals, Carlsbad, CA, 92008, USA
SO
     Nucleosides Nucleotides (1997), 16(7-9), 1625-1628
     CODEN: NUNUD5; ISSN: 0732-8311
     Marcel Dekker, Inc.
PB
DT
     Journal
LA
     English
AB
     Soln. phase prepn. of an oligodeoxyribonucleotide phosphorothioate
     (5'-TTGGGGTT) using phosphorothioate triester method is reported.
L11 ANSWER 27 OF 79 HCAPLUS COPYRIGHT 2000 ACS
     1997:741352 HCAPLUS
AN
DN
     128:34952
     A new approach to the synthesis of oligonucleotides and their
ΤI
     phosphorothicate analogs in solution
ΑU
     Reese, Colin B.; Song, Quanlai
     Dep. Chem., King's College London, London, WC2R 2LS, UK
CS
     Bioorg. Med. Chem. Lett. (1997), 7(21), 2787-2792
SO
     CODEN: BMCLE8; ISSN: 0960-894X
PB
     Elsevier Science Ltd.
DT
     Journal
LA
     English
AB
     A new approach to the synthesis of oligonucleotides and oligonucleotide
     phosphorothioates in soln. is described; it is based on H-phosphonate
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coupling at -40.degree. C, followed by in situ sulfur-transfer with

either

N-[(4-chlorophenyl)sulfanyl]phthalimide 19 or 4-[(2-cyanoethyl)sulfanyl]morpholine-3,5-dione 21.

- L11 ANSWER 28 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1997:669802 HCAPLUS
- DN 127:293567
- TI Chemical synthesis of peptides and polypeptides
- AU Sadat-Aalaee, Dean
- CS Biomeasure, Inc., Milford, MA, 01757, USA
- SO Protein-Based Mater. (1997), 3-35. Editor(s): McGrath, Kevin; Kaplan, David. Publisher: Birkhaeuser, Boston, Mass.

 CODEN: 65ECAZ
- DT Conference; General Review
- LA English
- AB A review with 213 refs. Topics include activation, coupling, protection and deprotection, as well as both soln. and solid-phase methods.
- L11 ANSWER 29 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1997:553998 HCAPLUS
- DN 127:234585
- TI Comparison of solution-phase and solid-phase syntheses of a restrained proline-containing analog of the nodularin macrocycle
- AU Webster, Kerri L.; Rutherford, Trevor J.; Gani, David
- CS Sch. Chem. and Centre Biomolecular Sciences, University, St.

Andrews/Fife,

KY16 9ST, UK

- SO Tetrahedron Lett. (1997), 38(32), 5713-5716 CODEN: TELEAY; ISSN: 0040-4039
- PB Elsevier
- DT Journal
- LA English
- AB The soln.-phase synthesis of a restrained (2S)-proline-contg. analog of the nodularin macrocycle,
- cyclo-[.beta.-ala-(2R)-Glu(.alpha.-OMe)-.gamma.-(2S)-Pro-(2R)-Asp(.alpha.-OMe)-.beta.-(2S)-Phe-], is described and compared to two solid-phase syntheses of the same cyclic isopentapeptide diester; one in which Fmoc-(2S)-Phe-.beta.-Ala-(2R)-Glu(.alpha.-OMe)-.gamma.-(2S)-Pro-(2R)-Asp(.alpha.-O-Wang Resin)-.beta.-OAllyl is deprotected and then cyclized on the resin and one in which this same precursor is removed from the resin prior to cyclization.
- L11 ANSWER 30 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1997:381350 HCAPLUS
- DN 127:81775
- ${\tt TI}$ Fluorinated peptides incorporating a 4-fluoroproline residue as potential inhibitors of HIV protease
- AU Tran, Thanh Thu; Patino, Nadia; Condom, Roger; Frogier, Tea; Guedj, Roger
- CS Lab. Chimie Bio-Organique, CNRS ERA 6001, Univ. Nice-Sophia Antipolis, Nice, 06108, Fr.
- SO J. Fluorine Chem. (1997), 82(2), 125-130 CODEN: JFLCAR; ISSN: 0022-1139
- PB Elsevier
- DT Journal
- LA English
- OS CASREACT 127:81775

AB Protected 4-fluoro-L-proline ester Fmoc-Pro(F)-OMe (I; Fmoc = 9-fluorenylmethoxycarbonyl) was prepd. as an attractive synthon for both solid and soln. phase peptide synthesis. Its use for the synthesis of Fmoc-Phe-Pro(F)-OMe and Fmoc-Pro(F)-Val-Val-OMe is presented. Direct fluorination with DAST of a 4-hydroxy proline residue incorporated into a peptide and elongation from the terminal amino group allowed prepn. of the hexapeptide Boc-Ala-Ala-Phe-Pro(F)-Val-Val-OMe, analogous to the p17-p24 gag junction of structural HIV proteins. None of the fluoropeptides in the paper displayed anti-protease or anti-HIV activity.

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L11 ANSWER 31 OF 79 HCAPLUS COPYRIGHT 2000 ACS
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1997:374851 HCAPLUS AN

DN 126:343816

ΤI Method for solution phase synthesis of oligonucleotides

Pieken, Wolfgang; Mcgee, Danny; Settle, Alecia; Zhai, Yansheng; Huang, IN

PA Nexstar Pharmaceuticals, Inc., USA; Pieken, Wolfgang; Mcgee, Danny; Settle, Alecia; Zhai, Yansheng; Huang, Jianping

PCT Int. Appl., 106 pp. SO

CODEN: PIXXD2

DTPatent

LA English LA

FAN.CNT 2																		
	PATENT NO.				KIND DATE					APPLICATION NO.					DATE			
ΡI	WO	9714706			Δ1 19970424				WO 1996-US16668					19961017				
	"							BR, BY, CA,							DIZ			
		w:																
															ΚZ,			
			LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	ΝZ,	PL,	PT,	RO,	RU,	SD,
			SE,	SG,	SI,	SK,	ТJ,	TM,	TR,	TT,	UA,	UG,	US,	UZ,	VN,	AM,	AZ,	BY,
			KG,	ΚZ,	MD,	RU,	ТJ,	TM										
		RW:	ΚE,	LS,	MW,	SD,	SZ,	UG,	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR,	GB,	GR,
			IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA		
	CA	2234159			AA 19970424				CA 1996-2234159 19961017									
	ΑU	9674518			A1 19970507				AU 1996-74518 19961017									
	ΑU	712779		B2 19991118														
	ΕP	863910			A1 19980916				E	P 19	96-93	3664	7	19961017				
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	IE, FI																	
	JP 2000500740			T2 20000125				JP 1997-516005					19961017					
	US	US 6001966			Α	A 19991214				US 1998-130232					19980806			
PRAI	US 1995-5619			9	19	19951019				•								

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WO 1996-US16668 19961017
     US 1997-780517 19970108
os
     MARPAT 126:343816
AB
     This invention discloses an improved method called PASS (product anchored
     sequential synthesis) for the soln. phase
     prepn. of oligodeoxyribonucleotides. The method PASS lends
     itself to automation and is ideally suited for large scale manuf. of
     oligodeoxyribonucleotides with high efficiency.
L11
     ANSWER 32 OF 79 HCAPLUS COPYRIGHT 2000 ACS
     1997:240708 HCAPLUS
AN
DN
     126:225558
ΤI
     Solution synthesis of peripheral acting analgesic opioid tetrapeptides
ΙN
     Rinaldi, Nicholas
PA
     Biochem Pharma Inc., Can.; Rinaldi, Nicholas
SO
     PCT Int. Appl., 26 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
     PATENT NO.
                        KIND DATE
                                             APPLICATION NO. DATE
                              -----
                                                -----
                                          WO 1996-CA552 19960815
     WO 9707129 A1 19970227
PΙ
          W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, AM, AZ,
              BY, KG, KZ, MD, RU, TJ, TM
          RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM
     AU 9666539
                         A1 19970312
                                              AU 1996-66539
                                                                   19960815
PRAI GB 1995-16994
                         19950818
     WO 1996-CA552
                        19960815
OS
     MARPAT 126:225558
AB
     This invention provides a bulk scale process for the soln. synthesis of
     enantiomerically pure, peripherally acting analgesic opioid tetrapeptides
     H-Tyr-R1-R2-R3-NH2, where R1 is D-Ala or D-Arg; R2 = R3 = Phe or
     p-fluorophenylalanine. The new and unique multi-step process includes coupling of X-Tyr-R1-OH (X= amino protecting group such as Boc) with
     H-R2-R3-NH2 using an activating agent such as N-hydroxysuccinimide, a
     neutralizing agent such as DIEA (diisopropylethylamine), and a suitable
     solvent such as DMF to yield the protected tetrapeptide. In this std.
     soln. phase synthesis, adjusting the
     individual factors (e.g., solvents, activating agents, neutralizing
     etc.) can minimize racemization of the second amino acid. Tremendous
cost
     efficiencies are achieved due to elimination of traditional sequential
     blocking-deblocking cycles and multiple chromatog. purifn. steps, such
     that these simple kilogram quantity methods can be scaled up to com.
     prodn.
L11
     ANSWER 33 OF 79 HCAPLUS COPYRIGHT 2000 ACS
     1997:188989 HCAPLUS
AN
DN
     126:277755
     Synthesis of [1,2-13C2] Gly and [2,2-2H2] Gly glutathione
TI
     Lu, Xiao-Ming; Fischman, Alan J.; Kenneway, Michael; Tompkins, Ronald G.;
ΑU
                     Searched by John Dantzman 703-308-4488
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Young, Vernon R.

- Surgical Service and Nuclear Medicine Division, Massachusetts General CS Hospital and Harvard Medical School, Boston, MA, 02114, USA
- J. Labelled Compd. Radiopharm. (1997), 39(3), 205-213 SO CODEN: JLCRD4; ISSN: 0362-4803
- PB Wiley
- DT Journal
- LA English
- AB [1,2-13C2] Gly and [2,2-2H2] Gly isotopomers of the intracellular tripeptide glutathione were prepd. by std. methods of soln. phase peptide synthesis. The synthetic products were characterized by gas chromatog./mass spectroscopy

(GC/MS) and 1H NMR spectroscopy. Optical purity was detd. by hydrolysis, derivatization of the free amino acids with isopropanol-acetyl chloride and pentafluoropropionic anhydride and NCI/MS with a Chirasil-Val

Heliflex

column. These compds. should serve as useful tracers for the non-invasive

study of glutathione synthesis and turnover rates in humans by GC/MS.

- ANSWER 34 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1997:15656 HCAPLUS
- DN 126:144528
- Design of specific structures using .alpha.,.beta.-dehydrophenylalanine ΤI residues: synthesis, crystal structure, and molecular conformation of Boc-L-Val-.DELTA.Phe-.DELTA.Phe-L-Val-.DELTA.Phe-.DELTA.Phe-L-Val-OCH3, a 310-helical heptapeptide
- AU
- Mitra, Shome Nath; Dey, Sharmistha; Karthikeyan, S.; Singh, Tej P. Department Biophysics, All India Institute Medical Sciences, New Delhi, CS 110029, India
- SO Biopolymers (1997), 41(1), 97-105 CODEN: BIPMAA; ISSN: 0006-3525
- PB Wiley
- Journal DT
- LA English
- AΒ To design an extensive 310-helical conformation, a heptapeptide Boc-L-Val-.DELTA.Phe-.DELTA.Phe-L-Val-.DELTA.Phe-.DELTA.Phe-L-Val-OCH3 (.DELTA.Phe = cis-.alpha.,.beta.-dehydrophenylalanine) with a repeat of two consecutive .DELTA. Phe residues has been synthesized using an azlactone method in soln. phase. The peptide was crystd. from its soln. in a methanol-water mixt. and its structure, where all peptide units are trans, has been detd. The peptide adopts a right-handed 310-helical conformation with more than two complete helical turns. Starting from

the

Boc group to the C-terminal residue of Val, the 310-helical structure is maintained well. The side chains of the four .DELTA. Phe residues in this helical arrangement exist in a slightly staggered arrangement. The solvent mol. (MeOH) forms two intermol. hydrogen bonds with the peptide, and thus, it helps to promote a head-to-tail packing of 310-helixes of

the

peptide. There are no lateral hydrogen bonds between the helixes, but there exist several van der Waals interactions involving the hydrophobic side chains of peptide mols.

- L11 ANSWER 35 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- 1996:682011 HCAPLUS AN
- DN 126:19172

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ΤI
     Soln. phase synthesis of
     oligodeoxyribonucleotide phosphorothioates
     Ravikumar, Vasulinga; Cole, Douglas L.
ΤN
     Isis Pharmaceuticals, Inc., USA
PA
SO
     U.S., 17 pp. Cont.-in-part of U.S. Ser. No. 99,075.
     CODEN: USXXAM
DT
     Patent
LA
     English
FAN.CNT 3
                       KIND DATE
     PATENT NO.
                                            APPLICATION NO. DATE
                             -----
PΙ
     US 5571902
                       Α
                             19961105
                                             US 1994-249442
                                                               19940526
                             19970325
                                             US 1993-99075
     US 5614621
                       Α
                                                               19930729
                                             CA 1994-2167671
     CA 2167671
                       AA
                             19950209
                                                               19940720
                                             WO 1995-US6825
                                                               19950526
                           19951207
     WO 9532980
                       A1
         W: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD, MG, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ,
             TM, TT
         RW: KE, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT,
             LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE,
             SN, TD, TG
     AU 9526570
                        Α1
                             19951221
                                             AU 1995-26570
                                                               19950526
                                            EP 1995-921510
     EP 766688
                        A1
                             19970409
                                                               19950526
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT,
SE
     US 6001982
                             19991214
                                             US 1996-692909
                                                               19960731
                        Α
     US 5847106
                        Α
                             19981208
                                             US 1997-789443
                                                               19970127
     US 6124450
                        Α
                             20000926
                                             US 1998-123138
                                                               19980727
PRAI US 1993-99075
                       19930729
     US 1994-249442
                       19940526
     WO 1995-US6825
                       19950526
     US 1997-789443
                       19970127
AB
     Soln. phase synthesis of
     oligodeoxyribonucleotide phosphorothioates is reported. Thus,
     oligodeoxyribonucleotide 5'-HO-TT dimer was prepd. via coupling of
     3'-acetylthymidine with thymidine phosphoramidite.
     ANSWER 36 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     1996:635221 HCAPLUS
AN
DN
     125:276590
     Solution phase synthesis of immunoregulating
ΤI
     peptides
     Deigin, Vladislav Isakovich; Korotkov, Andrei Marxovich
IN
PA
     Russia
     PCT Int. Appl., 10 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     Russian
FAN.CNT 1
                                             APPLICATION NO.
     PATENT NO.
                       KIND
                             DATE
                                                               DATE
                                             ______
                             _____
                                            WO 1996-RU46 19960228
     WO 9626955
                      A1 19960906
PΙ
         W: AU, BR, BY, CA, CN, CZ, HU, JP, KG, KP, KZ, LT, LV, MN, SK, UA,
             US, UZ
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                        C1 19980327
                                            RU 1995-102461
     RU 2107691
                                                               19950302
                    Searched by John Dantzman 703-308-4488
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CA 2214410
                       AA
                            19960906
                                           CA 1996-2214410 19960228
     AU 9649594
                       A1
                            19960918
                                           AU 1996-49594
                                                             19960228
     AU 708084
                       B2
                            19990729
     EP 818462
                       A1
                            19980114
                                           EP 1996-906117
                                                            19960228
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
ΙE
     CN 1185160
                                                            19960228
                       Α
                            19980617
                                           CN 1996-192317
     JP 11505515
                       T2
                            19990521
                                           JP 1996-526185
                                                            19960228
     BR 9607901
                                                             19960228
                      Α
                            19990908
                                           BR 1996-7901
                                                             19971002
     LV 11993
                      В
                            19980520
                                           LV 1997-186
                      В
     LT 4393
                            19981026
                                           LT 1997-158
                                                            19971002
                                           US 1998-894963
     US 6051683
                      Α
                            20000418
                                                            19980817
PRAI RU 1995-102461
                      19950302
     WO 1996-RU46
                      19960228
     CASREACT 125:276590; MARPAT 125:276590
     The invention relates to medicine, specifically, to method of obtaining
AΒ
     biol. active substances with immuno-regulating properties, and can be
     in medicine and veterinary science and in exptl. biochem.
fundamental
     problem addressed by the invention is that of producing a novel synthetic
     biol. active peptide with immuno-regulating properties and of the
     X-Glu-Trp-Y, in which X is H or Gly, Ala, Leu, Ile, Val, NVal, Pro, Tyr,
     Phe, Trp, D-Ala, D-Leu, D-Ile, D-Val, DNVal, D-Pro, D-Tyr, D-Phe, D-Trp,
     .gamma.-aminobutyric acid, .zeta.-aminocaproic acid; Y is Gly, Ala, Leu,
     Ile, Val, NVal, Pro, Tyr, Phe, Trp, D-Ala, D-Leu, D-Ile, D-Val, D-Pro, D-Tyr, D-Phe, D-Trp, .gamma.-aminobutyric acid,
.zeta.-aminocaproic
     acid, -OH, mono- or di-substituted amide (C1-C3). Peptide synthesis
     place in soln. by successive growth of a chain from the C terminus, using
     a strategy of max. blocking of functional groups, starting from amino
acid
     alkyl ester, using the method of activating the esters and the method of
     mixed anhydrides, using Boc amino acids. Thus, e.g., coupling of Boc-Ile
     pentafluorophenyl ester with Glu-Trp followed by deprotection with formic
     acid afforded H-Ile-Glu-Trp-OH (I) which was evaluated in the lymphocyte
     E-rosette formation assay in guinea pigs: E-rosette formation increased
     from 36.1% (after treatment with trypsin alone) to 61.4% (trypsin + 10-6
     mg/mL I) vs. 60.3% (trypsin + 10-6 mg/mL thymosin fraction 5).
L11 ANSWER 37 OF 79 HCAPLUS COPYRIGHT 2000 ACS
     1996:572036 HCAPLUS
AN
     125:222456
DN
ΤI
     Solution phase synthesis of blood platelet
     aggregation-inhibitory N-orotylpeptide and its intermediate peptides
     Okazaki, Takeo; Myazaki, Hiroshi
ΙN
     Shinnippon Seitetsu Kk, Japan; Shinnittetsu Kagaku
PΑ
     Jpn. Kokai Tokkyo Koho, 7 pp.
SO
     CODEN: JKXXAF
DT
     Patent
LA
     Japanese
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
     JP 08183797
                     A2 19960716
                                          JP 1994-327548
                                                            19941228
                   Searched by John Dantzman 703-308-4488
```

- AB The title peptide, orotyl-Ser-Arg-Gly-Asp-Trp-OH, which is a safe and potent blood platelet aggregation inhibitor (no data), was prepd. in a large scale by the soln. phase method involving sequential Boc-deprotection and coupling of Boc-Asp-(OBzl)-OH, Boc-Gly-OH, Boc-Arg(Z)2-OH, Boc-Ser(Bzl)-OH, and orotic acid to Boc-Trp(Z)-OBzl, and deprotection of Bzl and Z groups from the resulting orotyl-Ser(Bzl)-Arg(Z)2-Gly-Asp(OBzl)-Trp(Z)-OBzl.
- L11 ANSWER 38 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1996:529876 HCAPLUS
- DN 125:276443
- TI A solution-phase strategy for the parallel synthesis of chemical libraries
 - containing small organic molecules: a general dipeptide mimetic and a flexible general template
- AU Tarby, Christine M.; Cheng, Soan; Boger, Dale L.
- CS CombiChem, Inc., San Diego, CA, 92121, USA
- SO Mol. Diversity Comb. Chem.: Libr. Drug Discovery, Conf. (1996), 81-98. Editor(s): Chaiken, Irwin M.; Janda, Kim D. Publisher: American Chemical Society, Washington, D. C. CODEN: 63HMAW
- DT Conference; General Review
- LA English

in

- AB A general approach to the soln. phase, parallel synthesis of chem. libraries, which allows the prepn. of multi-milligram quantities of each individual member, is exemplified with both a dipeptide mimetic and flexible general template and is discussed
- this review, with 87 refs. In each step of the sequence, the reactants, unreacted starting material, reagents and their byproducts are removed by simple liq./liq. or liq./solid extns. providing the desired intermediates and final compds. in high purities (.gtoreq.90-100%) independent of the reaction yields and without deliberate reaction optimization.
- L11 ANSWER 39 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1996:339996 HCAPLUS
- DN 125:115115
- TI Synthesis of fragments of the peptide component of pseudobactin
- AU Okonya, John F.; Kolasa, Teodozyj; Miller, Marvin J.
- CS Department Chemistry Biochemistry, University Notre Dame, Notre Dame, IN, USA
- SO J. Pept. Sci. (1996), 2(3), 157-164 CODEN: JPSIEI; ISSN: 1075-2617
- DT Journal
- LA English
- AB Pseudobactin is a structurally complex and physiol. important siderophore (microbial iron chelator) from Pseudomonas putida-fluorescens. Various fragments of the unusual peptide component of pseudobactin were prepd. by soln.-phase peptide
 - synthesis. A class of related peptides named pseudomycins have shown promising antifungal activity. To examine if these peptide fragments above would elicit similar activity, the fragments were tested and found to have no antifungal activity in limited bioassays.
- L11 ANSWER 40 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1996:285951 HCAPLUS
- DN 125:34108

- TI Practical synthesis of disulfated hirudin C-terminal related peptides
- AU Okayama, Toru; Hongo, Tomoko; Nukui, Eriko; Muramatsu, Ryo; Hayashi, Hideya; Morikawa, Tadanori
- CS Research Laboratory, Fuji Chemical Industries, Ltd., Toyama, Japan
- SO Pept. Chem. (1996), Volume Date 1995, 33rd, 129-132 CODEN: PECHDP; ISSN: 0388-3698
- DT Journal
- LA English
- AB A symposium report on an improved practical procedure for the synthesis of
 - disulfated hirudin C-terminal related **peptides** by a **soln**. **phase synthesis** followed by a chem. O-sulfation of the tyrosine residues. In the course of this work, the authors obsd. an extensive racemization of the C-terminal amino acid residue in the O-sulfation process with pyridine-SO3 complex in a DMF-pyridine mixt.

The

authors found the reaction proceeds faster in the pyridine-free solvent system and the racemization of C-terminus was also suppressed; the desired

peptides were obtained in high yield.

- L11 ANSWER 41 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1996:285933 HCAPLUS
- DN 125:34100
- TI A solution-phase synthesis of fragment peptide derivatives using an automated synthesis apparatus
- AU Sugawara, Tohru; Kobayashi, Kyoko; Tanaka, Toshimasa; Fukushi, Shigeha; Kitada, Chieko; Fujino, Masahiko
- CS Molecular Chemistry Laboratory, Takeda Chemical Industries Ltd., Osaka, 532, Japan
- SO Pept. Chem. (1996), Volume Date 1995, 33rd, 57-60 CODEN: PECHDP; ISSN: 0388-3698
- DT Journal
- LA English
- ${\tt AB}$ A symposium report on the development of fully automated synthesis systems

for prepg. and isolating various kinds of pharmaceutical compds. As one application of the author's automated synthesis systems, a library of all possible dipeptides (25), tripeptides (125) and some tetrapeptide derivs. was synthesized systematically using 5 protected amino acids. The measured mol. optical rotation values for the library of 125 tripeptides correlate with the calcd. values obtained by summation of the mol. rotation values of the constituent amino acids. The app. has also been applied to the automated synthesis of 10 fragment tripeptides that are constituents of the hormone PACAP-27, and the soln.-phase synthesis of other tripeptide derivs. using

combinations of 10 different protected amino acids.

- L11 ANSWER 42 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1996:221847 HCAPLUS
- TI Synthesis of .gamma.-benzyl-.alpha.,L-glutamate oligomers and their star derivatives
- AU Wang, Xiaolan; Daly, William H.
- CS Department Chemistry, Louisiana State University, Baton Rouge, LA, 70803, USA
- SO Book of Abstracts, 211th ACS National Meeting, New Orleans, LA, March 24-28 (1996), POLY-112 Publisher: American Chemical Society, Washington, Searched by John Dantzman 703-308-4488

D. C.

CODEN: 62PIAJ

- DT Conference; Meeting Abstract
- LA English
- AB Highly monodisperse .gamma.-benzyl-.alpha.,L-glutamate oligomers (DP=4,8,12,16) have been synthesized by soln.

 phase convergent peptide synthesis. These peptides will be used to make model star polymers by coupling them to central units. Among the coupling methods studied, it is found that O-(7-azabenzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate(HATU) is the most effective coupling reagent for assembling BLG 4-mers. Efforts to couple BLG 8-mers and 16-mers are in process.
- L11 ANSWER 43 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1996:112931 HCAPLUS
- DN 124:290228
- TI Solution phase synthesis of Arg-Arg contained oligopeptides and studies on its activity
- AU Zhao, MIng; Peng, Shiqi; Wang, Yinye
- CS Coll. Pharmaceutical Sci., Beijing Med. Univ., Beijing, 100083, Peop.

Rep.

- China
- SO Zhongguo Yaowu Huaxue Zazhi (1995), 5(2), 91-5 CODEN: ZYHZEF
- DT Journal
- LA Chinese
- AB Oligopeptides Leu-Arg-Arg and Ser-Leu-Arg-Arg were prepd. by the soln. method, their vasodilation effect and inhibiting effect on ADP-induced platelet aggregation were obsd. The results indicated there was no differences between them and Arg-Arg dipeptide for vasodilation potency and their antiplatelet aggregating effect was also significant.
- L11 ANSWER 44 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1996:92221 HCAPLUS
- DN 124:261686
- TI Generalized Dipeptidomimetic Template: Solution Phase Parallel Synthesis of Combinatorial Libraries
- AU Boger, Dale L.; Tarby, Christine M.; Myers, Peter L.; Caporale, Lynn Helena
- CS Scripps Research Institute, La Jolla, CA, 92037, USA
- SO J. Am. Chem. Soc. (1996), 118(8), 2109-10 CODEN: JACSAT; ISSN: 0002-7863
- DT Journal
- LA English

GI

AB A simple and general approach to the **soln. phase,**Searched by John Dantzman 703-308-4488

the prepn. of multimilligram quantities of each individual member is described. In each step of the sequence, the reactants, unreacted starting material, reagents and their byproducts are removed by simple liq./liq. or liq./solid extn. providing the desired intermediates and final compds. in high purities (.gtoreq.90-95%) irresp. of reaction yields

and without deliberate reaction optimization.

- L11 ANSWER 45 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1995:982954 HCAPLUS
- DN 124:117945
- TI Quinazoline Antifolate Thymidylate Synthase Inhibitors: .gamma.-Linked L-D, D-D, and D-L Dipeptide Analogs of 2-Desamino-2-methyl-N10-propargyl-5,8-dideazafolic Acid (ICI 198583)
- AU Bavetsias, Vassilios; Jackman, Ann L.; Kimbell, Rosemary; Gibson, William;
 - Boyle, F. Thomas; Bisset, Graham M. F.
- CS CRC Centre for Cancer Therapeutics, Institute of Cancer Research, Sutton/ Surrey, SM2 5NG, UK
- SO J. Med. Chem. (1996), 39(1), 73-85 CODEN: JMCMAR; ISSN: 0022-2623
- DT Journal
- LA English
- AB The syntheses of .gamma.-linked L-D, D-D, and D-L dipeptide analogs of 2-desamino-2-methyl-N10-propargyl-5,8-dideazafolic acid (ICI 198583) are described. The general methodol. for the synthesis of these mols. involved the prepn. of the dipeptide derivs. employing soln. phase peptide synthesis

followed by condensation of the dipeptide free bases with the appropriate pteroic acid analog via di-Et cyanophosphoridate (DEPC) activation. In the final step, tert-Bu esters were removed by trifluoroacetic acid hydrolysis. Z-L-Glu-OBut-.gamma.-D-Ala-OBut, for example, was prepd.

from

.alpha.-tert-Bu N-(benzyloxycarbonyl)-L-glutamate and tert-Bu D-alaninate via isobutyl-mixed anhydride coupling. The Z-group was removed by catalytic hydrogenolysis and the resulting dipeptide free base condensed with 2-desamino-2-methyl-N10-propargyl-5,8-dideazapteroic acid via DEPC coupling. Finally, tert-Bu esters were removed by TFA hydrolysis to give ICI 198583-.gamma.-D-Ala. The compds. were tested as inhibitors of thymidylate synthase and L1210 cell growth. Good enzyme and growth inhibitory activity were found with .gamma.-linked L-D dipeptides, the best examples being the Glu-.gamma.-D-Glu deriv.(Ki = 0.19 nM, L1210 IC50 = 0.20 .+-. 0.017 .mu.M) and the Glu-.gamma.-D-.alpha.-aminoadipate deriv.

(Ki = 0.12 nM, L1210 IC50 = 0.13 .+-. 0.063 .mu.M). In addn., ICI 198583 L-.gamma.-D-linked dipeptides were resistant to enzymic degrdn. in mice.

- L11 ANSWER 46 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1995:834145 HCAPLUS
- DN 124:30383
- TI Application of a unique automated synthesis system for solution-phase peptide synthesis
- AU Sugawara, Tohru; Kobayashi, Kyoko; Okamoto, Shigeha; Kitada, Chieko; Searched by John Dantzman 703-308-4488

Fujino, Masahiko

- CS Mol. Chem. Lab., Pharmaceutical Res. Div., Osaka, 532, Japan
- SO Chem. Pharm. Bull. (1995), 43(8), 1272-80 CODEN: CPBTAL; ISSN: 0009-2363
- DT Journal
- LA English
- AB An automated synthesis system, which is suitable for repetitive syntheses using similar reaction procedures, was used to synthesize systematically

library of all possible dipeptides (25) and tripeptides (125) from 5 protected amino acids. The app. has also been applied to the automated synthesis of 10 fragment tripeptide derivs. that are constituents of the hormone PACAP-27. The measured mol. optical rotation values of the library of 125 tripeptides were found to correlate well with calcd.

values

а

obtained by summation of the mol. optical rotation values for the constituent amino acids.

- L11 ANSWER 47 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1995:631089 HCAPLUS
- DN 123:286627
- TI Peptide analogs of DNA consisting of L-.alpha.-amino-.gamma.-thymine butyric acid and L-valine subunits
- AU Ceulemans, G.; Khan, K.; Van Schepdael, A.; Herdewijn, P.
- CS Rega Inst. for Medical Res., Katholieke Univ. Leuven, Louvain, B-3000, Belg.
- SO Nucleosides Nucleotides (1995), 14(3-5), 813-16 CODEN: NUNUD5; ISSN: 0732-8311
- DT Journal
- LA English
- AB Reaction of N-Boc-L-homoserine benzylester with N3-benzoylthymine under Mitsumobu conditions afforded N-Boc-L-.alpha.-amino-.gamma.-N3-benzoylthymine butyric acid benzyl ester. After removal of the N-benzoyl and O-benzyl protecting group, this compd. was used in soln. phase peptide synthesis.
- L11 ANSWER 48 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1995:624512 HCAPLUS
- DN 123:314463
- TI Rapid solution phase synthesis of peptides by the Fmoc strategy
- AU Ueki, Masaaki; Tsurusaki, Takeshi; Okumura, Jin
- CS Department Applied Chemistry, Science University Tokyo, Tokyo, 162, Japan
- SO Pept. Chem. (1995), Volume Date 1994, 32nd, 213-16 CODEN: PECHDP; ISSN: 0388-3698
- DT Journal
- LA English
- AB New procedures for one-pot deprotection and coupling of peptides by the Fmoc strategy were developed.
- L11 ANSWER 49 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1995:549152 HCAPLUS
- DN 123:170059
- TI **Solution-phase synthesis** of phosphorothioate oligodeoxyribonucleosides by the phosphotriester method
- AU Barber, Isabelle; Imbach, Jean-Louis; Rayner, Bernard
- CS Laboratoire Chimie Bio-organique, Universite Montpellier II, Montpellier, Searched by John Dantzman 703-308-4488

34095, Fr.

SO Antisense Res. Dev. (1995), 5(1), 39-47

CODEN: AREDEI; ISSN: 1050-5261

DT Journal

LA English

AB A "phosphorothioate triester method" was investigated for the soln
.-phase synthesis of phosphorothioate
oligoribonucleosides. Using fully protected 3'-phosphorothiolate
thymidine bearing O-cyanoethyl and S-2,4-dichlorobenzyl groups as
phosphorothioate protecting groups, decathymidine nonaphorphorothioate

was

efficiently assembled through a blockwise procedure. Two side reactions occurred during the deprotection steps: breakage of inter-nucleoside linkages (1.8% per linkage) and formation of phosphate diester linkages (0.9%). Substitution of the dichlorobenzyl group by the more labile 4-nitrobenzyl S-protecting group reduced the extent of internucleoside bond breakage by one-half.

L11 ANSWER 50 OF 79 HCAPLUS COPYRIGHT 2000 ACS

AN 1995:495195 HCAPLUS

DN 122:291514

TI Silicon-Containing Amino Acids and Peptides. Asymmetric Synthesis of (Trialkylsilyl)alanines

AU Walkup, Robert D.; Cole, Derek C.; Whittlesey, Bruce R.

CS Department of Chemistry and Biochemistry, Texas Tech University, Lubbock, TX, 79409, USA

SO J. Org. Chem. (1995), 60(8), 2630-4 CODEN: JOCEAH; ISSN: 0022-3263

DT Journal

LA English

GΙ

The three (trialkylsilyl)alanines L-RCH2CH(NH2)CO2H (I; R = Me3Si, PhSiMe2, MeSiPh2) were synthesized in 6-9 steps from the com. available starting materials Me3SiCH2CH2CO2Na, ClSiMe2Ph, and ClSiPh2Me in 42%,

and 10% overall yields, resp., using an asym. .alpha.-bromination of the chiral N-acyloxazolidinone derivs. of the 3-(trialkylsilyl)propanoates to introduce the abs. configuration of the .alpha. center. Azide displacement, oxazolidinone removal, and redn. yielded I, which were converted to their N-(9-fluorenylmethoxycarbonyl) (Fmoc) derivs. for use in peptide synthesis. An x-ray crystal structure of azido(trimethylsilylpropanoyl)oxazolidinone II, an intermediate in the synthesis of I (R = SiMe3), substantiated the stereochem. course of the synthetic route. To demonstrate the ability of trialkylsilylalanines to undergo typical reactions assocd. with soln. phase

peptide synthesis in good yields, Fmoc-protected I (R = SiMe3) was coupled using DCC conditions to H-Phe-OMe, deprotected using diethylamine, coupled to Boc-Phe-OH (Boc = Me3CO2C), then deprotected using trifluoroacetic acid. The results reported indicate that amino acids bearing a variety of trialkylsilyl groups as large hydrophobic side chains can be synthesized by a general asym. synthesis route and incorporated into peptides.

- L11 ANSWER 51 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- 1995:455581 HCAPLUS AN
- ΤI New amino-protecting group, 2-adamantyloxycarbonyl (2-Adoc) and its application tot he synthesis of protected peptides
- ΑU Muklherjee, Ashis K.; Agosta, William C.
- CS
- Rockefeller Univ., New York, NY, USA Chemtracts: Org. Chem. (1994), 7(6), 415-16 SO CODEN: CMOCEI; ISSN: 0895-4445
- DTJournal
- English LA
- AB Researchers developed a new side-chain protecting group, 2-adamantyloxycarbonyl (2-Adoc) with the primary objective of increasing the soly. of the peptide fragment in org. solvents and of increasing stability to the conditions during the synthesis of protected peptide fragments to be used in convergent solid-phase peptide synthesis. 2-Adoc is shown to be suitable for .epsilon.-amino protection of lysine in convergent solid-phase peptide synthesis in combination with N.alpha.-fluoren-9-ylmethoxycarbonyl (Fmoc) protection and
- trifluoroacetic

acid-labile (TFA-labile) solid support. Researchers showed the stability and susceptibility of H-Lys-(2-Adoc)-OH (Fig. 1) to various acids and bases and found that, other than methanesulfonic acid, std. deprotecting agents, wuch as trifluoromethanesulfonic acid and hydrofluoric acid, worked satisfactorily. They also showed that various Fmoc and tert-butoxycarbonyl (Boc)-protected Lys-(2-Adoc) derivs. can be prepd. with the help of std. reagents. 2-Adoc-protected peptides were also used for solid-phase synthesis in combination with N.alpha.-Fmoc protection

and

TFA-cleavable resin support and were shown to be stable during piperidine treatment and TFA cleavage. Moreover, the fragments contg. the 2-Adoc groups were easily sol. in DMF in sufficient concn. for their use in fragment condensation. Researchers also showed that 2-Adose group protection in soln.-phase peptide synthesis was stable during the synthesis, including the deprotection of Boc groups.

- L11 ANSWER 52 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1995:418702 HCAPLUS
- DN 122:315025
- Protected 5-fluoro-2'-deoxyuridine monophosphate for solution-TIphase synthesis of oligodeoxyribonucleotides
- Mazzei, Mauro; Grandi, Teresa; Balbi, Alessandro; Abramova, Tatiana V.; Damonte, Gianluca; Silvestro, Luigi
- CS Ist. Sci. Farm., Genoa, 16132, Italy
- Farmaco (1994), 49(12), 793-7 SO CODEN: FRMCE8
- DT Journal
- LAEnglish
- os CASREACT 122:315025

AB In order to obtain new building blocks for oligodeoxyribonucleotide (ODN) soln. synthesis we are describing the synthesis of the protected dinucleotide I carrying 5-fluorouracil and thymine from 5-fluoro-2'-deoxyuridine as an example of future development in this field. I is in turn hydrolyzed to yield the unprotected dimer. The latter product could be esp. useful in the delivery of 5-fluorouracil from

L11 ANSWER 53 OF 79 HCAPLUS COPYRIGHT 2000 ACS

AN 1995:224860 HCAPLUS

DN 122:133795

TI Amino acids and peptides. Part 38. Development of a new amino-protecting group, 2-adamantyloxycarbonyl, and its application to peptide synthesis

AU Nishiyama, Yasuhiro; Shintomi, Noriyuki; Kondo, Yukihiro; Okada, Yoshio

CS Faculty Pharmaceutical Sciences, Kobe-Gakuin University, Kobe, 651-21, Japan

SO J. Chem. Soc., Perkin Trans. 1 (1994), (21), 3201-7 CODEN: JCPRB4; ISSN: 0300-922X

DT Journal

LA English

AB A new lysine .epsilon.-amino protecting group, 2-adamantyloxycarbonyl (2-Adoc), was developed, and its application to the solid-phase synthesis of protected peptides was demonstrated in combination with N2-fluoren-9-ylmethoxycarbonyl (Fmoc) protection and trifluoroacetic acid (TFA)-cleavable resin support. The 2-Adoc group was applied successfully also to the soln.-phase peptide synthesis depending on tert-butoxycarbonyl (Boc)-chem.

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ANSWER 54 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
AN
     1995:217625 HCAPLUS
DN
     122:133642
     2-Diphenylmethylsilylethyl (DPSE): a versatile protecting group for
ΤI
     oligodeoxyribonucleotide synthesis
     Ravikumar, Vasulinga T.; Cole, Douglas L.
ΑU
CS
     Isis Pharmaceuticals, Carlsbad, CA, 92008, USA
SO
     Gene (1994), 149(1), 157-61
     CODEN: GENED6; ISSN: 0378-1119
DΤ
     Journal
LA
     English
     2-Diphenylmethylsilylethyl (DPSE) is a new protecting group for the
AB
     internucleotidic bonds in the solid-support and soln. -
     phase synthesis of oligodeoxyribonucleotides
     by the phosphoramidite approach. This group is stable under acidic
     conditions and can be removed by a .beta.-fragmentation mechanism under
     mild conditions using aq. NH4OH. Alternatively, this group can also be
     removed using tetrafluorosilane in acetonitrile. Antiviral activity of
     oligodeoxyribonucleotide is reported (no data).
L11
     ANSWER 55 OF 79 HCAPLUS COPYRIGHT 2000 ACS
AN
     1994:509681 HCAPLUS
DN
     121:109681
     Liquid phase synthesis of peptides and peptide derivatives
TΙ
IN
     Sivruk, Gary A.; Eynon, John S.
PA
SO
     PCT Int. Appl., 28 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
                      KIND
     PATENT NO.
                            DATE
                                           APPLICATION NO.
                                                            DATE
PΙ
     WO 9325571
                      A1
                            19931223
                                           WO 1993-US5783
                                                            19930616
        W: AU, CA, JP, NZ, US
     EP 598899
                       A1
                            19940601
                                           EP 1993-916581
                                                            19930616
                            19980930
     EP 598899
                       В1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT,
SE
     JP 06509821
                       T2
                            19941102
                                           JP 1993-501810
                                                            19930616
     AU 671660
                       B2
                            19960905
                                           AU 1993-46381
                                                            19930616
     AT 171708
                       Ε
                            19981015
                                           AT 1993-916581
                                                            19930616
     ES 2123059
                      Т3
                            19990101
                                           ES 1993-916581
                                                            19930616
     US 5516891
                      Α
                            19960514
                                           US 1994-190111
                                                            19940525
                      19920616
PRAI IE 1992-1942
     WO 1993-US5783
                      19930616
AB
     A continuous liq. phase peptide synthesis method for prepg. peptides
     contg. 2-10 amino acid residues uses (1) Fmoc as the protecting group for
     the non-side chain amino functionality, (2) NH3 or a primary or secondary
     amine to remove the Fmoc protecting group, and (3) a substituted
     carbodiimide as the coupling agent in a proper org. solvent. Thus,
     H-Pro-OtBu.HCl and Fmoc-Lys(BOC)-OH were stirred 2 h with
     diisopropylcarbodiimide and Et3N in CH2Cl2. The resulting suspension was
     treated with 4-aminomethylpiperidine and stirred for 1 h followed by
     filtration and washing of the filtrate with pH 5.5 phosphate buffer.
     soln. was dried over Na2SO4, filtered, concd., treated with
     Fmoc-Asp(OtBu)-OH and diisopropylcarbodiimide, and stirred 1 h.
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Deprotection, workup, coupling with Fmoc-Ser(OtBu)-OH, and deprotection were carried out as before; the tetrapeptide soln. was then treated with Ac2O and Et3N to give a solid which was stirred with CF3CO2H to give 65% Ac-Ser-Asp-Lys-Pro-OH.

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L11 ANSWER 56 OF 79 HCAPLUS COPYRIGHT 2000 ACS
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AN 1994:292660 HCAPLUS

DN 120:292660

- TI The synthesis and use of pp60src-related peptides and phosphopeptides as substrates for enzymic phosphorylation studies
- AU Perich, John W.; Meggio, Flavio; Valerio, Robert M.; Johns, R. B.; Pinna, Lorenzo A.; Reynolds, Eric C.
- CS Sch. Chem., Univ. Melbourne, Parkville, 3052, Australia
- SO Bioorg. Med. Chem. (1993), 1(5), 381-8 CODEN: BMECEP; ISSN: 0968-0896
- DT Journal
- LA English
- AB A series of peptides and phosphopeptides corresponding to the auto-phosphorylation site of pp60src, -Asn-Glu-Tyr416-Thr-Ala-, were prepd. by either Boc/soln. or Fmoc/solid phase peptide synthesis and used as substrates to study their enzymic phosphorylation by various casein kinases. The Tyr(P)-contg. peptide, Asn-Glu-Tyr(P)-Thr-Ala, was prepd.

by
the use of Fmoc-Tyr(PO3Bzl2)-OH in Fmoc/solid phase peptide synthesis
followed by acidolytic treatment of the peptide-resin with 5%
anisole/CF3CO2H. Both Asn-Glu-Tyr-Thr-Ala and Asn-Glu-Ser(P)-Thr-Ala
were

prepd. by the Boc/soln. phase peptide synthesis and employed hydrogenolytic deprotection of the protected peptides. Enzymic phosphorylation studied established that (A) the Tyr residue acted as an unusual pos. determinant for directing phosphorylation to the Thr-residue, (B) the rate of Thr-phosphorylation was markedly facilitated by a change from the Tyr-residue to the Tyr(P)-residue, and (C) a Ser(P)-residue was as effective as the Tyr(P)-residue in facilitating Thr-phosphorylation. A subsequent structure-function study using Asn-Glu-Phe-Thr-Ala, Asn-Glu-Tyr(Me)-Thr-Ala (prepd. by Fmoc/solid phase peptide synthesis) and

Asn-Glu-Cha-Thr-Ala (prepd. by hydrogenation of Asn-Glu-Tyr-Thr-Ala) established that the rate

of Thr-phosphorylation was influenced by the extent of hydrophobic-hydrophobic interactions by the aralkyl side-chain group (either arom. or aliph.) of the 416-residue with casein kinase-2; the rate

of Thr-phosphorylation being decreased by the introduction of Me or hydroxyl groups at the 4-position of the arom. group {i.e. Tyr(Me) and Tyr

resp.) but enhanced by the introduction of the hydrophilic phosphate group

{i.e. as Tyr(P)}.

- L11 ANSWER 57 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1993:650272 HCAPLUS
- DN 119:250272
- TI Scale-up of oligonucleotide synthesis. Solution phase
- AU Seliger, H.
- CS Polym. Sect., Univ. Ulm, Ulm, Germany
 Searched by John Dantzman 703-308-4488

SO Methods Mol. Biol. (Totowa, N. J.) (1993), 20(Protocols for Oligonucleotides and Analogs), 391-435 CODEN: MMBIED; ISSN: 1064-3745

DT Journal; General Review

LA English

AB A review with 228 refs. on the soln. phase prepn. of oligodeoxyribonucleotides.

L11 ANSWER 58 OF 79 HCAPLUS COPYRIGHT 2000 ACS

AN 1993:192250 HCAPLUS

DN 118:192250

TI Solution-phase segment synthesis of boron-rich peptides

AU Kane, Robert R.; Pak, Roger H.; Hawthorne, M. Frederick

CS Dep. Chem. Biochem., Univ. California, Los Angeles, CA, 90024-1569, USA

SO J. Org. Chem. (1993), 58(5), 991-2 CODEN: JOCEAH; ISSN: 0022-3263

DT Journal

LA English

OS CASREACT 118:192250

GΙ

Fmoc
$$=$$
 Gly-NH $=$ CO $=$ OCMe3 $=$ (CH2)3 SiMe2CMe3 $=$ B10H10

AB Small peptides I (Fmoc = 9-fluorenylmethoxycarbonyl; n = 1, 2, 4), contg. up to 40 boron atoms, were efficiently synthesized in soln. Condensation of a closo-carborane amino ester with Fmoc-Gly-F afforded the orthogonally

protected dipeptide I (n=1) in good yield. Selective removal of protecting groups allowed segment condensations, culminating with prodn. of the octapeptide I (n=4). The lipophilic closo-carboranes in these peptides could be readily converted to their hydrophilic anionic nido derivs. This methodol. should find utility in the precise synthesis of boron-rich macromols., and should be esp. suited for use in the antibody mediated boron neutron capture therapy of cancer.

L11 ANSWER 59 OF 79 HCAPLUS COPYRIGHT 2000 ACS

AN 1993:125027 HCAPLUS

DN 118:125027

TI A new and simplified method for hydrogenolytic deprotection in solution-phase peptide synthesis

Ι

AU Pallenberg, Alexander J.

CS Procyte Corp., Kirkland, WA, 98034, USA

SO Tetrahedron Lett. (1992), 33(50), 7693-6

CODEN: TELEAY; ISSN: 0040-4039

```
Journal
DT
LA
     English
     An improved method for the deprotection of synthetic peptides by
AB
catalytic
     hydrogenation is described. The new method allows for precise control of
     counterion stoichiometry and affords the peptides in high purity and
     yield, while avoiding the problems usually assocd. with conventional
     deprotection methods.
     ANSWER 60 OF 79 HCAPLUS COPYRIGHT 2000 ACS
T.11
ΑN
     1993:125013 HCAPLUS
DN
     118:125013
ΤI
     Solution phase synthesis and conformational
     analysis of Glu-Ser-Leu-Ser-Ser-Ser-Glu-Glu-NHMe and its peptide
congeners
     (non-phosphorylated region 14-21 of bovine .beta.-casein A2)
ΑU
     Perich, John W.; Johns, R. B.
     Sch. Chem., Univ. Melbourne, Parkville, 3052, Australia
CS
     Aust. J. Chem. (1992), 45(11), 1857-69
SO
     CODEN: AJCHAS; ISSN: 0004-9425
DT
     Journal
     English
LΑ
AB
     The octapeptide H-Glu-Ser-Leu-Ser-Ser-Glu-Glu-NHMe.CF3CO2H and its
     five shorter peptide congeners (from tripeptide to heptapeptide) were
     prepd. in high yield and purity by the tert-butoxycarbonyl mode of
     soln. phase peptide synthesis
     followed by palladium-catalyzed hydrogenolytic deprotection of the six
     protected peptides in 50% CF3CO2H/CH3CO2H soln. The anal. of the six
     peptides by 13C NMR spectroscopy and C18 reversed-phase chromatog.
     suggested that a structural arrangement commenced at the hexapeptide
stage
     and was considered to be due to the formation of a .beta.-turn
     conformation.
     ANSWER 61 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     1993:60059 HCAPLUS
ΑN
DN
     118:60059
ΤI
     Benextramine-neuropeptide Y receptor interactions: contribution of the
     benzylic moieties to [3H]neuropeptide Y displacement activity
     Doughty, Michael B.; Chaurasia, Chandra S.; Li, Ke
ΑU
     Sch. Pharm., Univ. Kansas, Lawrence, KS, 66045-2506, USA
CS
     J. Med. Chem. (1993), 36(2), 272-9
CODEN: JMCMAR; ISSN: 0022-2623
SO
DT
     Journal
     English
LA
     CASREACT 118:60059
OS
     Benextramine (BXT) analogs [RCH2NH(CH2)6NHCH2CH2S]2.4HCl (I; R =
AB
     m-MeOC6H4,p-MeOC6H4, o-ClC6H4, m-ClC6H4, p-ClC6H4, 2-naphthyl, o-HOC6H4,
     m-HOC6H4, p-HOC6H4, H) were synthesized using soln.-
     phase peptide synthesis methodol. and analyzed
     for activity in displacing specifically bound 1nM N-[propionyl-
     3H]neuropeptide Y([3H]NPY) from benextramine-sensitive neuropeptide Y
     (NPY) binding sites in rat brain. The new synthetic approach to these
     analogs began with the acylation of cystamine with the
     N-hydroxysuccinimide ester of tert-butyloxycarbonyl (Boc) protected
     6-aminohexanoic acid, followed by deprotection of the Boc groups with 4N
     HCl in dioxane. Acylation of this sym. diammine with
N-hydroxysuccinimidSearched by John Dantzman
                                                703-308-4488
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esters of appropriately substituted benzoic acids, followed by redn. of the resultant tetramides with diborane in refluxing THF, afforded the target compds. The BXT analog lacking the benzylic group {i.e., I (R = H0} had no [3H] NPY displacement activity at concns. up to $1.4 \times 10^{-3} \text{ M}$. The 9-fold range in activities obsd. for the ortho, meta and para regioisomers of the methoxy, chloro, and hydroxy benextramine analogs at benextramine-sensitive NPY rat brain binding sites does not differ from the range of potencies obsd. at .alpha.-adrenoceptors. However, the

order

of potencies at at [3H]-NPY sites differs from the orders of potencies at .alpha.-adrenoceptors, with analogs I (R = m-MeOC6H4, m-HOC6H4, 2-naphyhyl) being the most active at [3H]-NPY binding sites. The present results demonstrate the importance of the benzylic moiety for BXT's NPY antagonist activity, and suggest that the BXT binding site on the NPY receptor is significantly distinct from that on the .alpha.-adrenoceptor.

- L11 ANSWER 62 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1993:22612 HCAPLUS
- DN 118:22612
- TI Efficient solution-phase synthesis of multiple O-phosphoseryl-containing peptides related to casein and statherin
- AU Perich, John W.; Kelly, David P.; Reynolds, Eric C.
- CS Sch. Dent. Sci., Univ. Melbourne, Melbourne, Australia
- SO Int. J. Pept. Protein Res. (1992), 40(2), 81-8 CODEN: IJPPC3; ISSN: 0367-8377
- DT Journal
- LA English
- AB The multiple phosphoserine-contg. peptides R-[Ser(PO3H2)]n-Glu-Glu-NHMe.cntdot.CF3CO2H (R = H, n = 3; R = H-Asp, H-Glu, n = 2) were prepd. using Boc-Ser(PO3Ph2)-OH (Boc = tert-butoxycarbonyl) in the Boc mode of soln. phase peptide synthesis

followed by Pt-mediated hydrogenolytic deprotection of the Ser(PO3Ph2)-contg. peptides. The protected peptides were assembled using the mixed anhydride coupling methods with 40% CF3CO2H/CH2Cl2 used for removal of the Boc group from intermediate Boc-protected peptides.

- L11 ANSWER 63 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1992:551398 HCAPLUS
- DN 117:151398
- TI Preparation of nonapeptides as gonodoliberin antagonists
- IN Koenig, Wolfgang; Sandow, Juergen; Kolar, Cenek
- PA Hoechst A.-G., Germany
- SO Eur. Pat. Appl., 16 pp.
- CODEN: EPXXDW
- DT Patent
- LA German
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
ΡI	EP 477499	A1	19920401	EP 1991-112817	19910730	
	EP 477499	B1	19940126			
	R: AT, BE,	CH, DE	, DK, ES, FR,	GB, GR, IT, LI, LU	, NL, SE	
	AT 100822	E	19940215	AT 1991-112817	19910730	
	ES 2062628	Т3	19941216	ES 1991-112817	19910730	
	NO 9103020	A	19920205	NO 1991-3020	19910802	
	CA 2048407	AA	19920205	CA 1991-2048407	19910802	
	Searched by John Dantzman 703-308-4488					

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AU 9181548
                        A1
                             19920206
                                             AU 1991-81548
                                                               19910802
     AU 641035
                        B2
                              19930909
     ZA 9106097
                        Α
                             19920429
                                             ZA 1991-6097
                                                               19910802
     IL 99062
                        A1
                             19950731
                                             IL 1991-99062
                                                               19910802
     JP 05148299
                        A2
                             19930615
                                             JP 1991-219139
                                                               19910805
     US 5434138
                        Α
                             19950718
                                             US 1993-151056
                                                               19931112
PRAI DE 1990-4024779
                       19900804
     EP 1991-112817
                       19910730
                       19910801
     US 1991-739233
OS
     MARPAT 117:151398
     Peptides X-A-B-C-Ser-D-E-F-G-Pro-H [I; X = C2-8 alkanoyl; A =
AB
     D-3-(2-naphthyl)alaninyl (D-Nal), D-Phe, D-Trp all of which may be
     substituted on the arom. ring; B = (substituted) D-Phe; C = D-3-(3-pyridyl)alaninyl (D-Pal), (substituted) D-Phe, -D-Trp; D = Tyr, His; E = D-Ser(R1); R1 = glycosyl group; F = Leu, Trp, Phe; G = Ser(R1);
     H, Gly-NH2, D-Ala-NH2, azaGly-NH2] were prepd. as gonadoliberin
     antagonists which inhibit testosterone and estrogen biosynthesis.
     Ac-D-Nal-D-p-Cl-Phe-D-Pal-Ser-Tyr-D-Ser(Rha)-Leu-Ser(Rha)-Pro-D-Ala-NH2
     (II) (Rha = rhamnosyl) was prepd. via std. soln.
     phase peptide synthesis starting from
     Fmoc-Pro-OH and H-D-Ala-NH2.HCl using the appropriate protected amino
     acids. II at 60 .mu.q/24 h via minipump infusion in rats inhibited
     testosterone synthesis.
     ANSWER 64 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     1992:236128 HCAPLUS
AN
DN
     116:236128
     Synthesis of the simple peptide model Ac-Abu(PO3H2)-NHMe
TI
     Valerio, Robert M.; Perich, John W.; Alewood, Paul F.; Tong, Glenn;
ΑU
Johns,
     Sch. Chem., Univ. Melbourne, Parkville, 3052, Australia
CS
ŞO
     Aust. J. Chem. (1992), 45(4), 777-84
     CODEN: AJCHAS; ISSN: 0004-9425
DT
     Journal
     English
LA
     The simple model substrate Ac-L-Abu(PO3H2)-NHMe(Abu(PO3H2) =
AΒ
     NHCH(CH2CH2PO3H2)CO] was prepd. by the use of the protected
     4-(diethylphosphono)butanoic acid deriv. Boc-Abu(PO3Et2)-OH (Boc =
     Me3CO2C) in the Boc mode of soln. phase
     peptide synthesis. The protected peptide model
     Ac-Abu(PO3Et2)-NHMe was prepd. by initial reaction of the
     isobutoxycarbonyl mixed anhydride of Boc-Abu(PO3Et2)-OH with MeNH2
     followed by cleavage of the Boc group from Boc-Abu(PO3Et2)-NHMe with 4 M
     HCl/dioxane and N-acetylation of H-Abu(PO3Et2)-NHMe.HCl with the
     isobutoxycarbonyl mixed anhydride of AcOH. Cleavage of the phosphonate
Εt
     groups was effected with 33% HBr/AcOH or 10% BrSiMe3/MeCN to give
     Ac-L-Abu(PO3H2)-NHMe in nearly quant. yield.
     ANSWER 65 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     1991:608523 HCAPLUS
ΑN
DN
     115:208523
TΙ
     Solution-phase synthesis of the potassium
     channel blocker, charybdotoxin
     Lambert, Paul F.; Kuroda, Hisaya; Chino, Naoyoshi; Watanabe, Takushi X.;
ΑU
     Kimura, Terutoshi; Sakakibara, Shumpei
                    Searched by John Dantzman 703-308-4488
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CS
     Protein Res. Found., Pept. Inst., Inc., Minoh, Japan
     Pept. 1990, Proc. Eur. Pept. Symp., 21st (1991), Meeting Date 1990, 111-12. Editor(s): Giralt, Ernest; Andreu, David. Publisher: ESCOM Sci.
SO
     Publ., Leiden, Neth.
     CODEN: 57HNAI
DT
     Conference
LA
     English
GI
pGlu-Phe-Thr-Asn-Val-Ser-Cys-Thr-Thr-Ser-Lys-
 Glu-Cys-Trp-Ser-Val-Cys-Gln-Arg-Leu-His-Asn-
 Thr-Ser-Arg-Gly-Lys-Cys-Met-Asn-Lys-Lys-Cys-
 Arg-Cys-Tyr-Ser-OH
                                                 Ι
AB
     A symposium report on the soln.-phase
     synthesis of charybdotoxin with peptide sequence I (pGlu
     = pyroglutamic acid). The linear peptide was oxidized to give the
     disulfide form. The disulfide bridges in the synthetic product were
found
     to be between Cys7-Cys28, Cys13-Cys33 and Cys17-Cys35.
L11 ANSWER 66 OF 79 HCAPLUS COPYRIGHT 2000 ACS
     1991:229369 HCAPLUS
AN
DN
     114:229369
ΤI
     Synthesis of casein-related peptides and phosphopeptides. IX. A
modified
     method for the synthesis of Ser(P) peptides by using Ppoc-Ser(PO3Bzl2)-OH
     Perich, John W.; Alewood, Paul F.; Johns, R. B.
ΑU
CS
     Sch. Chem., Univ. Melbourne, Parkville, 3052, Australia
SO
     Aust. J. Chem. (1991), 44(3), 377-87
     CODEN: AJCHAS; ISSN: 0004-9425
DT
     Journal
     English
LA
     Benzyl phosphate groups were sensitive to acid conditions, and a
AΒ
stability
     study with dibenzyl iso-Bu phosphate under various acid conditions is
     described. While extensive acidolytic debenzylation of the dibenzyl
     phosphorotriester Boc-Ser(PO3R2)-Leu-OR (I; Boc = Me3CO2C, R = CH2Ph)
     occurred on treatment with either 4 M HCl/dioxane or 50% CF3CO2H/CH2Cl2,
     only minor benzyl loss occurred with the use of HCO2H or 1 M HCl/AcOH.
     Minimization of benzyl phosphate loss during the synthesis of a dibenzyl
     phosphoserine-contg. tripeptide was effected by the use of 98% HCO2H (or
1
     M HCl/AcOH) for the cleavage of the Boc group from I. In alternative
     procedure, the protected 2-phenylisopropyloxycarbonyl deriv.
     Me2CPh02C-Ser(PO3R2)-OH (R = CH2Ph) was prepd. by an efficient four-step
     procedure and was used in a soln.-phase
     peptide synthesis for the high-yielding prepn.
     of Boc-Glu(OR)-Ser(PO3R2)-Leu-OR (R = OH2Ph). The protected tripeptide
     was deprotected by palladium-catalyzed hydrogenolysis in formic acid and
     gave H-Glu-SerPO3H2-Leu-OH in near quant. yield.
                   Searched by John Dantzman
                                                703-308-4488
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L11
    ANSWER 67 OF 79 HCAPLUS COPYRIGHT 2000 ACS
     1991:82523 HCAPLUS
AN
DN
     114:82523
TI
     An efficient facilitated method for solution phase
     peptide synthesis
     Head, David B.
ΑU
CS
     Lab. Rational Drug Design, Univ. Hosp., Boston, MA, 02118, USA
SO
     Pept.: Chem., Struct. Biol., Proc. Am. Pept. Symp., 11th (1990), Meeting
     Date 1989, 1012-14. Editor(s): Rivier, Jean E.; Marshall, Garland R.
     Publisher: ESCOM Sci. Pub., Leiden, Neth.
     CODEN: 56XTA7
DT
     Conference
LA
     English
AB
     A symposium report on the use of a cholestane moiety as a bulky cryst.
     handle for the soln.-phase synthesis of
     peptides.
    ANSWER 68 OF 79 HCAPLUS COPYRIGHT 2000 ACS
T.11
     1990:632007 HCAPLUS
ΑN
DN
     113:232007
TΙ
     Solution-phase synthesis of porcine brain
     natriuretic peptide (pBNP) using S-trimethylacetamidomethylcyste
     ine
     Kiso, Yoshiaki; Yoshida, Makoto; Kimura, Tooru; Fujiwara, Yoichi;
ΑU
     Shimokura, Masanori; Akaji, Kenichi
     Dep. Med. Chem., Kyoto Pharm. Univ., Kyoto, 607, Japan
CS
SO
     Chem. Pharm. Bull. (1990), 38(5), 1192-9
     CODEN: CPBTAL; ISSN: 0009-2363
DT
     Journal
LA
     English
     The hexadodecapeptide corresponding to the entire amino acid sequence of
AB
     porcine brain natriuretic peptide (pBNP) was synthesized by assembling
     four segments in soln., followed by HF deprotection and subsequent oxidn.
     to establish an intramol. disulfide bridge. The synthesis using the
newly
     developed S-trimethylacetamidomethylcysteine deriv. gave a better yield
     than that using the S-2,4,6-trimethylbenzylcysteine deriv. The chick
     rectum relaxant activity of the synthetic pBNP was 2.9 times more potent
     than that of .alpha.-rat atrial natriuretic peptide (.alpha.-rANP).
     ANSWER 69 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
     1990:612687 HCAPLUS
AN
DN
     113:212687
ΤI
     Preparation of tripeptides via solution phase coupling using
     propylphosphonic anhydride
     Flemming, Hans Wolfram; Rukwied, Manfred; Schmidt, Manfred
ΙN
PΑ
     Hoechst A.-G., Fed. Rep. Ger.
SO
     Ger. Offen., 4 pp.
     CODEN: GWXXBX
DT
     Patent
```

FAN.CNT 1					
PATENT NO.	KIND	DATE	APE	PLICATION NO.	DATE
PI DE 3839379	A1	19900523	DE	1988-3839379	19881122
CA 1335493	A1	19950509	CA	1989-614545	19890929
	Searched	by John Dantz	man	703-308-4488	

LΑ

German

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A2
                              19900530
                                              EP 1989-121277
                                                                19891117
     EP 370399
     EP 370399
                        A3
                              19910918
     EP 370399
                        В1
                              19950621
         R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE
     ES 2075028
                                              ES 1989-121277
                        Т3
                              19951001
                                                                19891117
     DK 8905844
                        Α
                              19900523
                                              DK 1989-5844
                                                                19891121
     AU 8945328
                        A1
                              19900531
                                              AU 1989-45328
                                                                19891121
     AU 626608
                        B2
                              19920806
                              19900903
     JP 02219587
                        A2
                                              JP 1989-300960
                                                                19891121
     JP 2843618
                        B2
                              19990106
     US 5191065
                        Α
                              19930302
                                              US 1991-728028
                                                                19910708
PRAI DE 1988-3839379
                       19881122
     US 1989-438073
                       19891120
     MARPAT 113:212687
OS
AΒ
     U-A-B-C-OH (U = H, urethane protecting group; A, B = naturally occurring
     .alpha.-amino acid residue or deriv.; C = arom. .alpha.-aminoacid
     residue), were prepd. by 1) reaction of U1-B-OH (U1 = hydrogenolyzable
     urethane protecting group) with H-C-OR (R = C1-4 alkyl) in the presence
of
     propylphosphonic anhydride (I), 2) hydrogenolysis of the coupling product
     to give H-B-C-OR, 3) coupling of the latter with U-A-OH in the presence
of
     I, and 4) enzymic cleavage of the R group. Thus, a mixt. of Z-Ser-OH,
     H-Tyr-OMe.HCl, NaCl, EtOAc, and N-ethylmorpholine at pH 5.0 was treated
     with I over 30 min at .ltoreq.30.degree.. The EtOAc phase was hydrogenolyzed over Pd/C with addn. of aq. HCl to maintain pH 4.0.
     aq. phase contq. the hydrogenolyzed dipeptide was coupled with Z-Trp-OH
as
     above and the product in H2O/EtOAc was stirred with trypsin at
     35-40.degree. for 7 h to give 42% Z-Trp-Ser-Tyr-OH of 98.2% purity.
     ANSWER 70 OF 79 HCAPLUS COPYRIGHT 2000 ACS
L11
AN
     1990:591863 HCAPLUS
DN
     113:191863
     In situ silylation with trimethylsilyl cyanide. An outstanding protocol
TΙ
     for fast peptide synthesis. A synopsis
     Anteunis, M. J. O.; Becu, C.; Becu, F.
ΑU
     Lab. Org. Chem., State Univ. Ghent, Ghent, B-9000, Belg. Bull. Soc. Chim. Belg. (1990), 99(6), 361-77
SO
```

CS

CODEN: BSCBAG; ISSN: 0037-9646

- DTJournal; General Review
- LA English
- The title protocol is discussed with 34 refs. The use of trimethylsilyl cyanide as a potent "in situ" silylating agent and its compatibility with AB most classical functionalities employed during soln.

phase peptide syntheses allows repetitive peptide chain elongations (including linear head-to-tail) with a min. of chem. steps and manipulations. The outstanding features are: the upscaling facilities, the simplicity and the high purity of the final peptides exempt of stereomutation.

- ANSWER 71 OF 79 HCAPLUS COPYRIGHT 2000 ACS L11
- 1990:235818 HCAPLUS AN
- DN 112:235818
- TISolution syntheses of two enkephalin-containing peptides, peptide E and dynorphin(1-24), using Nin-(2,4,6-triisopropylphenylsulfonyl)tryptophan
- Kitagawa, Kouki; Kawamoto, Tatsuhiko; Futaki, Shiroh; Kiyama, Shinya; ΑU Searched by John Dantzman 703-308-4488

Akita, Tadashi; Moritoki, Hideki; Kiso, Yoshiaki

- CS Fac. Pharm. Sci., Univ. Tokushima, Tokushima, 770, Japan
- SO Chem. Pharm. Bull. (1989), 37(10), 2631-8 CODEN: CPBTAL; ISSN: 0009-2363
- DT Journal
- LA English
- OS CASREACT 112:235818
- AB Two enkephalin-contg. peptides, peptide E and dynorphin (1-24), were synthesized by conventional soln. methods employing a new tryptophan deriv., Nin-(2,4,6-triisopropylphenylsulfonyl)tryptophan [H-Trp(Tps)-OH]. All protecting groups employed, including the Tps group, were removed by treatment with 1 M CF3SO3H-PhSMe inCF3CO2H at the final steps of these syntheses. Subsequent purifications by Sephadex G-25 chromatog., CM-Biogel A ion exchange chromatog., and reversed-phase HPLC afforded highly purified samples. Both synthetic peptide E and dynorphin (1-24) exhibited high in vitro opioid activity. The usefulness of this new tryptophan deriv. for practical peptide synthesis was established through these syntheses of complex tryptophan-contg. peptides.
- L11 ANSWER 72 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1990:179858 HCAPLUS
- DN 112:179858
- TI Synthesis of O-phosphotyrosine-containing peptides. II. Solution-phase synthesis of
- Asn-Glu-Ptyr-Thr-Ala through methyl phosphate protection
 AU Valerio, Robert M.; Perich, John W.; Kitas, Eric A.; Alewood, Paul F.;
- Johns, R. B.
 CS Dep. Org. Chem., Univ. Melbourne, Parkville, 3052, Australia
- SO Aust. J. Chem. (1989), 42(9), 1519-25 CODEN: AJCHAS; ISSN: 0004-9425
- DT Journal
- LA English
- OS CASREACT 112:179858
- AB The O-phosphotyrosine pentapeptide H-Asn-Glu-Tyr(PO3H2)-Thr-Ala-OH.CF3CO2H, which is a naturally occurring sequence from the autophosphorylated Rous sarcoma virus pp60v-src, was prepd. in high yield from Boc-Tyr(PO3Me2)-OH (Boc = Me3CO2C) by a soln.-phase method. The protected pentapeptide Z-Asn-Glu(OBzl)-Tyr(PO3Me2)-Thr(Bzl)-Ala-OBzl (Z = PhCH2O2C; Bzl = PhCH2) was deprotected by a two-stage procedure which involved initial Pd-catalyzed hydrogenolysis followed by the removal of the phosphate Me group with BrSiMi3/MeCN, BrSiMe3/PhSMe in CF3CO3H, or CF3SO3H/CF3COH/Me2S/m-cresol.
- L11 ANSWER 73 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- AN 1989:574652 HCAPLUS
- DN 111:174652
- TI Studies on **peptides**. CLXIV. **Solution-phase synthesis** of a 36-residue **peptide** amide corresponding to the entire amino acid sequence of chicken antral peptide
- AU Guo, Lili; Murayama, Eigoro; Funakoshi, Susumu; Fujii, Nobutaka; Aono, Mitsuru; Matsuda, Masayuki; Moriga, Motoyuki; Yajima, Haruaki
- CS Fac. Pharm. Sci., Kyoto Univ., Kyoto, 606, Japan
- SO Chem. Pharm. Bull. (1988), 36(11), 4364-76 CODEN: CPBTAL; ISSN: 0009-2363
- DT Journal
- LA English
- OS CASREACT 111:174652

GΙ

and

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H-Phe-Leu-Pro-His-Val-Phe-Ala-Glu-Leu-Ser-Asp-
  Arg-Lys-Gly-Phe-Val-Gln-Gly-Asn-Gly-Ala-Val-
  Glu-Ala-Leu-His-Asp-His-Phe-Tyr-Pro-Asp-Trp-
  Met-Asp-Phe-NH2
                                                  Ι
     A 36-residue peptide amide corresponding to the entire amino acid
sequence
     of chicken antral peptide (I) was synthesized by assembling seven peptide
     fragments via the azide, followed by PhSMe-mediated deprotection with Me3SiBr and Me3SiO3SCF3 in CF3CO2H. The synthetic peptide stimulated
     gastric secretion, but not pancreatic secretion.
    ANSWER 74 OF 79 HCAPLUS COPYRIGHT 2000 ACS
     1988:455220 HCAPLUS
ΑN
DN
     109:55220
     Applications of cobalt(III) complexes in solid and solution
TΙ
     phase peptide syntheses
ΑU
     Mensi, Nahla E.
     Rutgers, State Univ., New Brunswick, NJ, USA
CS
     (1987) 178 pp. Avail.: Univ. Microfilms Int., Order No. DA8723271
SO
     From: Diss. Abstr. Int. B 1988, 48(7), 1976
DT
     Dissertation
LA
     English
AB
     Unavailable
L11 ANSWER 75 OF 79 HCAPLUS COPYRIGHT 2000 ACS
     1987:554744 HCAPLUS
AN
     107:154744
DN
TΤ
     Synthesis of casein-related peptides and phosphopeptides. I.
     Solution-phase synthesis and carbon-13 NMR
     spectroscopy of the N-.alpha.-acetyl octapeptide N-methylamide
     corresponding to region 14-21 of bovine .beta.-casein A2
     Perich, John W.; Alewood, Paul F.; Johns, R. B.
ΑU
     Dep. Org. Chem., Univ. Melbourne, Parkville, 3052, Australia
CS
     Aust. J. Chem. (1987), 40(2), 257-71
SO
     CODEN: AJCHAS; ISSN: 0004-9425
DT
     Journal
LA
     English
OS
     CASREACT 107:154744
AB
     Title octapeptide Ac-Glu-Ser-Leu-Ser-Ser-Glu-Glu-NHMe (I) was
     synthesized by the soln.-phase method by using the mixed anhydride
     coupling procedure for the fragment condensation of
Ac-Glu(OBut)-Ser(But)-
     Leu-OH with H-Ser(But)-Ser(But)-Glu(OBzl)-Glu(OBzl)-NHMe.HCl, followed by
     palladium-catalyzed hydrogenolysis of Ac-Glu(OBut)-Ser(But)-Leu-Ser(But)-
     Ser(But)-Ser(But)-Glu(OBzl)-Glu(OBzl)-NHMe in trifluoroacetic acid. The
     synthesis of the two peptide fragments was accomplished in high yields
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purity by using the repetitive excess mixed anhydride procedure and the isobutoxycarbonyl mixed anhydride of acetic acid for the rapid and high

703-308-4488

Searched by John Dantzman

yielding N-acetylation of the tripeptide fragment. 13C NMR spectroscopy was routinely used to monitor the efficiency of the coupling steps and to confirm the structure of I, signal assignments being possible for both the protected tri- and pentapeptides. ANSWER 76 OF 79 HCAPLUS COPYRIGHT 2000 ACS L111987:214347 HCAPLUS AN DN 106:214347 TIProperties of Nin-(2,4,6-triisopropylphenylsulfonyl)tryptophan and its application to the synthesis of .delta.-sleep inducing peptide Kiso, Yoshiaki; Shimokura, Masanori; Narukami, Takatomo; Nakamura, AU Akihiro; Shiomi, Hirohito Kyoto Pharm. Univ., Kyoto, 607, Japan
Pept. Chem. (1986), Volume Date 1985, 23rd, 131-6 CS SO CODEN: PECHDP; ISSN: 0388-3698 DT Journal English LA The 2,4,6-triisopropylphenylsulfonyl (Tps) group was introduced into the AB indole ring of Z(OMe)-Trp-OCH2Ph [Z(OMe) = 4-MeOC6H4CH2O2C] by treatment with Tps-Cl under phase-transfer catalytic conditions to give ${\tt Z\,(OMe)\,\hbox{-}Trp\,(Tps)\,\hbox{-}OCH2Ph.}$ The Tps group was stable under acidic (CF3CO2H, CF3CO2H/thioanisole, 25% HBr/AcOH) and basic (1N NaOH, 80% N2H4) conditions but easily removed in CF3SO3H-thioanisole-CF3CO2H. Z(OMe)-Trp(Tps)-OH was used in the soln. phase synthesis of .delta.-sleep inducing peptide, H-Trp-Gly-Gly-Asp-Ala Ser-Gly-Glu-OH. ANSWER 77 OF 79 HCAPLUS COPYRIGHT 2000 ACS 1.11 ΑN 1986:460933 HCAPLUS DN 105:60933 ΤI Studies on peptides. CXXXVI. Solution-phase synthesis of a 37-residue peptide amide corresponding to the entire amino acid sequence of human calcitonin gene-related peptide Fujii, Nobutaka; Otaka, Akira; Funakoshi, Susumu; Nomizu, Motoyoshi; Akaji, Kenichi; Yajima, Haruaki; Yamamoto, Itsuo; Torizuka, Kanji; ΑU Kitagawa, Kouki; et al. Kyoto Univ., Kyoto, 606, Japan
Chem. Pharm. Bull. (1986), 34(2), 613-20 CS SO CODEN: CPBTAL; ISSN: 0009-2363 DT Journal LA English CASREACT 105:60933 OS GT

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H-Ala-Cys-Asp-Thr-Ala-Thr-Cys-Val-
Thr-His-Arg-Leu-Ala-Gly-Leu-Leu-
Ser-Arg-Ser-Gly-Gly-Val-Val-Lys-
Asn-Asn-Phe-Val-Pro-Thr-Asn-Val-
Gly-Ser-Lys-Ala-Phe-NH2
```

Ι

The title peptide (I) was prepd. by a series of azide fragment AB condensations in soln. from 7 protected peptide segments. The final protected 37-peptide amide was deblocked by CF3SO3H/thioanisole in CF3CO2H

and the resulting deblocked peptide was cyclized by air oxidn. to give I. The 1-adamantyl (Ad) group was used for the protection of the SH group of cysteine; the Ad group was cleaved by the above acidolysis or by (CF3CO2)T1.

- ANSWER 78 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- 1983:122481 HCAPLUS ΑN
- DN 98:122481
- ΤI Purification of synthetic analogs of yeast mating hormone by reversed-phase chromatography
- Shenbagamurthi, P.; Naider, Fred; Becker, Jeffrey M.; Steinfeld, Alvin S. Coll. Staten Island, City Univ. New York, Staten Island, NY, 10301, USA ΑU
- CS
- J. Chromatogr. (1983), 256(1), 117-25 CODEN: JOCRAM; ISSN: 0021-9673 SO
- DT Journal
- LA English
- The .alpha.-type cells of Saccharomyces cerevisiae secrete low-mol.-wt. AB peptides, termed .alpha.-factors, which affect the sexual conjugation between .alpha.- and a-mating types of this yeast. The tridecapeptide .alpha.-factor (Trp-His-Trp-Leu-Gln-Leu-Lys-Pro-Gly-Gln-Pro-Met-Tyr), the dodecapeptide .alpha.-factor
- (His-Trp-Leu-Gln-Leu-Lys-Pro-Gly-Gln-Pro-Met-
 - Tyr), and a series of 8 analogs, were synthesized without

purifn. of intermediates, using std. soln. phase

techniques of peptide synthesis. Crude peptides

(125-500 mg) were loaded on to a preparative .mu.Bondapak C18 column (Waters Prep LC/System 500) and eluted with MeOH-H2O-trifluoroacetic acid (TFA) mixts. The recovery of purified peptide was as high as 93%.

factor analogs had biol. activity similar to that of the natural peptides.

The incorporation of TFA (.ltoreq.0.025%) in the mobile phase provides excellent conditions for the sepn. and purifn. of peptides. TFA has a significant effect on both peak shape and retention time in the concn. range 0-0.25%.

- L11ANSWER 79 OF 79 HCAPLUS COPYRIGHT 2000 ACS
- 1976:560496 HCAPLUS ΑN
- DN 85:160496
- Combined peptide synthesis method using peptide formation on insoluble TΙ supports and in solutions
- Shvachkin, Yu. P.; Ryabtsev, M. N.; Zuyanova, T. I.; Funtova, S. M.; ΑU Ivanovskaya, L. V.; Levinskii, A. B.
- CS Inst. Eksp. Endokrinol. Khim. Gorm., Moscow, USSR
- Zh. Obshch. Khim. (1976), 46(3), 717 SO CODEN: ZOKHA4
- DTJournal
- LA Russian
- Me3CO2C-Pro-Lys(CO2CH2Ph)-Thr-OMe was prepd. by the title procedure. Key AB steps included successive condensation of excess Me3CO2C-Lys(CO2CH2Ph)-OC6H4NO2-4 (I) with polymer-bound threonine (II), filtration, and Searched by John Dantzman 703-308-4488 reaction

of the filtrate with addnl. II. Residual I was filtered and condensed with Thr-OMe in soln.

=> d 1-14 bib abs

ANSWER 1 OF 14 WPIDS COPYRIGHT 2000 L12 DERWENT INFORMATION LTD 2000-292826 [25] AN WPIDS N2000-219598 DNC C2000-088439 DNN ΤI New high molecular weight form of endostatin, useful e.g. as antiangiogenic agent for treating cancer, isolated from hemofiltrate of patients with kidney failure. A88 B04 D16 S03 DC FORSSMANN, W; STAENDKER, L IN (HAEM-N) HAEMOPEP PHARMA GMBH PA CYC WO 2000017240 A1 20000330 (200025)* DE PΙ RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE WO 2000017240 A1 WO 1999-EP6963 19990921 PRAI DE 1999-19926040 19990608; DE 1998-19842992 19980921; DE 1999-19915267 19990403 AN 2000-292826 [25] WO 200017240 A UPAB: 20000524 AB NOVELTY - High molecular weight endostatin (hE) produced from the hemofiltrate of patients with renal insufficiency. DETAILED DESCRIPTION - The patient's blood is hemofiltered through a cellulose triacetate filter of exclusion limit 20 kD, then the hemofiltrate acidified, cooled to 4 deg. C and chromatographed on a exchange column as described in J. Chromatogr., A, 776 (1997) 125. The individual eluate pools (pH pools) are fractionated on a reverse-phase C4 column, eluting with a gradient of 0-30% B to 7 min then 30-65% B to 77 min (A = 0.1 vol.% trifluoroacetic acid (TFA); B = 80 vol.% acetonitrile, 0.1 vol.% TFA). The eluate fractions are screened for hE by mass spectrometry. INDEPENDENT CLAIMS are also included for the following: (1) pharmaceutical composition containing hE; (2) antibodies (Ab) against hE or its synthetic fragments; (3) diagnostic agent containing Ab; (4) nucleic acid encoding hE; and (5) determining the concentration of hE in the blood; ACTIVITY - Antitumor; antiproliferative. MECHANISM OF ACTION - hE inhibits angiogenesis. USE - hE is used to treat; (i) diseases that involve uncontrolled angiogenesis, particularly tumors; and (ii) vascular diseases of supporting or connective tissue, respiratory tract, cardiovascular system, urogenital tract and nervous system, or sensory organs (particularly the eye). hE is also used to raise specific antibodies which are used for diagnosis and treatment of conditions that involve overexpression of hE. ADVANTAGE - hE has a very long plasma half-life and can be administered repeatedly without inducing an immune response. Dwq.0/0

Searched by John Dantzman 703-308-4488

DERWENT INFORMATION LTD

L12 ANSWER 2 OF 14 WPIDS COPYRIGHT 2000

WPIDS

2000-246195 [21]

DNC C2000-074483

AN

```
New benzamidine compounds are platelet aggregation inhibitors for
treating
     e.g. thrombosis, stroke, myocardial infarction, inflammation,
     arteriosclerosis and metastasis.
DC
     B02 B03
IN
     BOVY, P R; RICO, J G; ROGERS, T E
PA
     (SEAR) SEARLE & CO G D
CYC
    1
PΙ
    US 6037365
                  A 20000314 (200021)*
                                              15p
ADT US 6037365 A US 1998-160089 19980925
PRAI US 1998-160089
                      19980925
AN
     2000-246195 [21]
                        WPIDS
          6037365 A UPAB: 20000502
AB
     NOVELTY - Benzamidine compounds (I) are new.
          DETAILED DESCRIPTION - Benzamidine compounds of formula (I) and
their
     salts are new.
          R1, R2 = H, halo, alkoxy, alkyl or hydroxy;
          W' = H, alkyl, alkenyl, aryl or alkoxycarbonyl (all optionally
     substituted by alkyl or aryl (optionally substituted by halo, alkoxy or
     alkyl));
          A = alkyl, alkenyl, alkynyl or alicyclyl (all optionally
substituted
    by OH, alkoxy, alkyl, halo or aryl (optionally substituted by halo, NO2,
     alkoxy or alkyl));
          Z' = a group of formula (i) or (ii);
          R3, R4 = H, halo, alkoxy, alkyl, sulfonyl, arylsulfonyl,
     heterocyclyl, phenyl (optionally substituted by halo, alkoxy or alkyl),
or
    phosphate, phosphinate or phosphonate (attached via P and optionally
    O-substituted by one or more alkyl, aryl, alkenyl or H);
    u = 1 \text{ or } 2;
       = 0-2;
          Q = one or more H, halo, OH, alkyl or alkoxy; and
          R9 = H, halo, carboxyl, alkoxycarbonyl, alkyl or alkoxy.
          ACTIVITY - Antiaggregant; Thrombolytic; Cerebroprotective; Cardiant;
     Antiinflammatory; Antiarteriosclerotic; Cytostatic.
          In assays 3S-((4-((4-(aminoiminomethyl)phenyl)amino)-1,4-
     dioxobutyl)amino)-4-hydroxy-(4-fluorophenyl)butanoic acid
trifluoroacetate
     had an IC50 value for platelet aggregation in canine platelet rich plasma
     in vitro of 0.15 (no units are given).
          MECHANISM OF ACTION - None given
          USE - As platelet aggregation inhibitors (claimed) for treating e.g.
     thrombosis, stroke, myocardial infarction, inflammation, arteriosclerosis
     and metastasis.
     Dwg.0/0
    ANSWER 3 OF 14 WPIDS COPYRIGHT 2000
                                            DERWENT INFORMATION LTD
L12
     2000-205976 [18]
                        WPIDS
AN
DNC
     C2000-063689
     New heptapeptide luteinizing hormone releasing hormone analogs used to
TΙ
     modulate levels of sex hormones and used in the treatment of e.g. benign
     prostate hypertrophyl, prostate tumors, breast and ovary tumors etc...
DC
     DWIGHT, W J; GREER, J; HAVIV, F; NICHOLS, C J
ΙN
     (ABBO) ABBOTT LAB
PΑ
                   Searched by John Dantzman
                                               703-308-4488
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CYC
     WO 2000009544 A1 20000224 (200018) * EN
PΤ
        RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE
         W: CA JP MX
     WO 2000009544 A1 WO 1999-US17874 19990806
PRAI US 1999-232425
                      19990115; US 1998-133055
                                                 19980812
     2000-205976 [18]
                        WPIDS
AB
     WO 200009544 A UPAB: 20000412
     NOVELTY - Heptapeptide luteinizing hormone releasing hormone (LHRH)
     analogs of formula (I) and their salts, esters and prodrugs them are
          DETAILED DESCRIPTION - Heptapeptide LHRH analogs of formula (I) and
     their salts, esters and prodrugs thereof are new: R1-A-B-C-D-E-F-G-R2
(I).
          R1 = lower alkylcarbonyl;
          A = 3-(2-naphthyl)-D-alanyl, (3-(4-chloro))-D-phenylalanyl or
     sarcosyl;
          B = 3-(1-naphthyl)-D-alanyl or (3-(4-chloro))-D-phenylalanyl;
          C = 3-(3-pyridyl)-D-alanyl or 3-(1-naphthyl)-D-alanyl;
     D = seryl;
          E = arginyl, (N-epsilon-nicotinyl)lysyl, N-methylphenylalanyl,
     (4-(3-amino-1,2,4-triazol-5-yl))phenylalanyl,
(4-(3-amino-1, 2, 3-triazol-5-
     yl))-N-methylphenylalanyl, (4-(N-acetyl))-N-methylphenylalanyl,
     (4-(N-nitro))-N-methylphenylalanyl, (4-(N-acetyl))-phenylalanyl, tyrosyl,
     N-methyltyrosyl or 1,2,3,4-tetrahydroisoquinoline-3-carbonyl;
          F = D-arginyl, D-asparaginyl, D-citrulluyl, D-glutamyl,
     D-homocitrully1, D-2-amino-6-NG, NG-diethylguanidinohexanoy1,
     (N-epsilon-nicotinyl)-D-lysyl, (4-(3-amino-1,2,4-triazol-5-yl))-D-
     phenylalanyl, (4-(N-acetyl))-D-phenylalanyl or D-tryptyl;
          G = cyclohexylalanyl, leucyl or N-methylleucyl;
     R2 = NR4 R5;
          R4 = H, Me or Et;
          R5 = lower alkyl or lower alkyl-R6;
          R6 = NH2, guanidino, H, OH, phenyl, morpholinyl, piperidinyl,
     pyrrolyl, pyridyl, pyrrolidinyl, pyrrolidinonyl or quinuclidinyl wherein
     the piperidinyl, pyrrolyl, pyrrolidinyl and pyrrolidinonyl are optionally
     substituted by a methyl group.
          INDEPENDENT CLAIMS are also included for the following:
          (1) a pharmaceutical formulation comprising (I); and
          (2) a method of modulating gonadotropin hormones in a mammal
     comprising administering (I).
          ACTIVITY - Cytostatic; gynocological; analgesic; depilatory.
          MECHANISM OF ACTION - Modulator of sex hormone levels, and (I) have
     activity as LHRH agonists or antagonists
          USE - (I) can be used to modulate the levels of gonadotropin and
     androgen secretion in male and female mammals. They can be used to treat
     conditions such as benign prostate hypertrophy, dysmenorrhea,
     endometriosis, precocious puberty, prostate cancer, uterine fibrosis,
     prostate necrosis, breast and ovary tumors, cryptorchidism, hirsutism,
     gastric motility disorders and other sex hormone dependent diseases.
     Dwg.0/0
L12 ANSWER 4 OF 14 WPIDS COPYRIGHT 2000 DERWENT INFORMATION LTD
     2000-038633 [03]
                        WPIDS
AN
DNC
     C2000-009856
     Liquid phase carriers for synthesis
                   Searched by John Dantzman 703-308-4488
```

```
of biopolymers in solution, particularly of proteins and nucleic acids.
DC
     B04 D16
IN
     KOESTER, H; WOERL, R
PA
     (KOES-I) KOESTER H
CYC
     84
PΙ
     WO 9955718
                   A2 19991104 (200003)* EN
                                              87p
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL
            OA PT SD SE SL SZ UG ZW
         W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB
            GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LR LS LT LU LV
            MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT
            UA UG UZ VN YU ZA ZW
                   A 19991116 (200015)
     AU 9936643
ADT WO 9955718 A2 WO 1999-US8939 19990426; AU 9936643 A AU 1999-36643
19990426
FDT AU 9936643 A Based on WO 9955718
PRAI US 1998-67337
                      19980427
     2000-038633 [03]
                        WPIDS
AN
          9955718 A UPAB: 20000118
AΒ
     NOVELTY - Liquid phase carrier (LPC) comprises a polyvalent group (Sp)
     with more than two points of attachment that carry reactive groups (X1)
     for synthesis of biopolymers.
          DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the
     following:
          (a) sequential solution phase synthesis
     of biopolymers on LPC; and
          (b) LPC coupled to biopolymers.
          ACTIVITY - None given.
          MECHANISM OF ACTION - None given.
          USE - LPC are used for solution-phase
     synthesis of peptides, peptide nucleic acids,
     oligosaccharides and particularly oligonucleotides, especially for
     therapeutic applications.
          ADVANTAGE - Solution-phase synthesis on
     LPC can provide (kilo) gram scale quantities of biopolymers, with high
     purity and better yields than possible with known solution methods. LPC,
     and its reaction products formed during biopolymer synthesis, are soluble
     in the reaction medium and can be modified to have other advantageous
     properties such as compatibility with chromatography. The considerable
     difference in size between products and reagents makes possible
     purification by gel-permeation chromatography and products can be
analyzed
     by mass spectrometry (of the fully protected material), allowing direct
     monitoring of the synthesis process.
     Dwq.0/0
L12 ANSWER 5 OF 14 WPIDS COPYRIGHT 2000
                                            DERWENT INFORMATION LTD
     1999-590442 [50]
                        WPIDS
ΑN
DNC C1999-172356
     Isolated protein used as a laxative in the treatment of constipation.
ΤI
DC
     CURRIE, M G; FOK, K F; WIEGAND, R C
IN
PA
     (SEAR) SEARLE & CO G D
CYC
    1
     US 5969097
                  A 19991019 (199950)*
PΙ
                                              14p
ADT US 5969097 A US 1992-903029 19920623
PRAI US 1992-903029
                     19920623
                   Searched by John Dantzman 703-308-4488
```

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AN
     1999-590442 [50]
                        WPIDS
          5969097 A UPAB: 19991201
AB
     NOVELTY - An isolated protein containing a 15 amino acid sequence as
given
     in the specification, is new.
          DETAILED DESCRIPTION - An isolated protein containing a 15 amino
acid
     sequence of formula (I) (human guanylin) is new.
          Pro-Gly-Thr-Cys-Glu-Ile-Cys-Ala-Tyr-Ala-Ala-Cys-Thr-Gly-Cys
                                                                         (I).
          An INDEPENDENT CLAIM is also included for an isolated protein
     consisting of (I).
    ACTIVITY - Laxative.
          MECHANISM OF ACTION - Intestinal quanylate cyclase regulator.
          The figure shows the bioactivity of human guanylin in the T84 cell
     bioassay. Comparison of the activity of human guanylin with rat guanylin
     indicates that they have similar potency to activate intestinal guanylate
     cyclase. Both types of guanylin are about one order of magnitude less
     potent than STs, which are heat stable enterotoxins that activate
     intestinal guanylate cyclase.
          USE - The protein can be used as a laxative in the treatment of
     constipation.
          DESCRIPTION OF DRAWING(S) - The figure shows the bioactivity of
human
     quanylin in the T84 cell bioassay.
     Dwg. 3a/7
    ANSWER 6 OF 14 WPIDS COPYRIGHT 2000
                                            DERWENT INFORMATION LTD
     1999-570511 [48]
AN
                        WPIDS
DNC
    C1999-166443
     Nalpha-2(p-biphenyl)-propyloxycarbonyl amino acid pentafluoro-phenyl
ΤI
     esters used in syntheses of polypeptide chains, peptides and proteins.
DC
     A96 A97 B04 B05
     CAREY, R I
IN
     (UYGE-N) UNIV GEORGIA RES FOUND INC
PΑ
CYC
     US 5952497
                  A 19990914 (199948)*
PΙ
                                              16p
ADT US 5952497 A Provisional US 1996-21499 19960710, US 1997-891676 19970710
PRAI US 1996-21499
                      19960710; US 1997-891676 19970710
     1999-570511 [48]
ΑN
                        WPIDS
          5952497 A UPAB: 19991122
AΒ
     US
     NOVELTY - N alpha -2(p-biphenyl)-propyloxycarbonyl amino acid
     pentafluoro-phenyl esters.
          DETAILED DESCRIPTION - N alpha -2(p-biphenyl)-propyloxycarbonyl
amino
     acid pentafluoro-phenyl esters are of formula Bpoc-Xxx-Pfp.
          Xxx = amino acid excluding esters in which amino acid is
     S-(acetamidomethyl)-L-cysteine or L-(tertiary butyl)-glutamic acid;
          Bpoc = 2-(p-biphenylyl)propyloxycarbonyl; and
          Pfp = pentafluorophenyl.
          INDEPENDENT CLAIMS are also included for:
          (1) 3,4-dihydro-4-oxo-1,2,3-benzotriazin-3-yl (ODhbt) esters of N
     alpha -2(p-biphenyl)propyloxycarbonyl amino acids;
          (2) compounds of formula (I); and
          (3) compounds of formula (II).
          R and R' = H, alkyl, optionally substituted cycloalkyl, optionally
     substituted aryl.
                   Searched by John Dantzman 703-308-4488
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(I) is not N alpha -2(biphenyl)-propyloxycarbonyl-L-glutamine
     pentafluorophenyl ester, N alpha
-2 (biphenyl) -propyloxycaronyl-L-glutamate
     pentafluorophenyl ester or N alpha -2(biphenyl)-propyloxycarbonyl-S-
     acetamidomethyl)-L-cysteine pentafluorophenyl ester.
          USE - Used in syntheses of polypeptide chains (claimed) as well as
     peptides and proteins.
          ADVANTAGE - Used to improve syntheses of polypeptide chains
     (claimed). Are storage-stable crystalline materials or storage-stable
     amorphous solids. Facilitate and simplify both solid- and solution
     -phase peptide synthesis especially in
     automated peptide synthesizers by eliminating need for
     activations, filtrations and couplings prior to peptide bond-forming
     reaction. Purification of peptides prepared in solution is facilitated by
     substantial lack of by-products. Can be used in combination with resin
     linkages not stable to repetitive basic reagents used to remove N alpha
     -Fmoc groups. Used in combination with side-chain protecting groups and
     resin linkages removable with trifluoroacetic acid/scavenger mixtures,
     distinguishing them from analogous N alpha -Boc derivatives that requires
     side-chain protecting groups and resin linkages removable only with
     stronger acid/scavenger mixtures. Facilitate peptide synthesis with N
     alpha -Bpoc amino acids compared with prior art N alpha -Bpoc amino acid
     cyclohexylamine or dicyclohexylamine salts that require tedious
     manipulation to activate the storage stable salts for peptide couplings.
     Facilitate peptide synthesis compared with other N alpha -Bpoc amino acid
     active esters whose reactivity is too sluggish to be useful in practical
     application to solid-phase peptide synthesis.
     Dwg.0/0
    ANSWER 7 OF 14 WPIDS COPYRIGHT 2000
                                            DERWENT INFORMATION LTD
L12
     1998-399055 [34]
AN
                        WPIDS
CR
     1997-258645 [23]
DNC
    C1998-120896
TΙ
     Solution phase synthesis of
     oligonucleotide(s) and peptide(s) - useful for large scale
     automated preparation of oligonucleotide(s) and peptide(s).
DC
IN
     GOLD, L; PIEKEN, W
PΑ
     (NEXS-N) NEXSTAR PHARM INC; (PROL-N) PROLIGO LLC
CYC
    82
     WO 9830578
                   A1 19980716 (199834)* EN 103p
PΙ
        RW: AT BE CH DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA
            PT SD SE SZ UG ZW
        W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE
            GH GM GW HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG
            MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG
            UZ VN YU ZW
     AU 9860223
                   Α
                     19980803 (199850)
     US 5874532
                   Α
                     19990223 (199915)
     US 6001966
                   Α
                      19991214 (200005)
                   A1 20000503 (200026)
     EP 996627
                                        EN
         R: AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE
    WO 9830578 A1 WO 1998-US562 19980106; AU 9860223 A AU 1998-60223
19980106;
     US 5874532 A US 1997-780517 19970108; US 6001966 A Div ex US 1994-289654
     19940812, CIP of WO 1996-US16668 19961017, Cont of US 1997-780517
     19970108, US 1998-130232 19980806; EP 996627 A1 EP 1998-903457 19980106,
                   Searched by John Dantzman
                                             703-308-4488
```

WO 1998-US562 19980106

FDT AU 9860223 A Based on WO 9830578; US 6001966 A Cont of US 5874532; EP 996627 A1 Based on WO 9830578

PRAI US 1997-780517 19970108; US 1994-289654 19940812; WO 1996-US16668 19961017; US 1998-130232 19980806

AN 1998-399055 [34] WPIDS

CR 1997-258645 [23]

AB WO 9830578 A UPAB: 20000531

Solution phase synthesis of peptides

comprises:

- (a) reacting an N-terminal protected amino acid monomer unit with a peptide starting material to form a reaction mixture containing a peptide product, and
- (b) partitioning the peptide product from the unreacted peptide starting material, unreacted N-terminal protected amino acid monomer unit,

side-products and reagents based on the presence of the N-terminal protecting group.

The product of the reaction is also claimed.

Also claimed is a method for the solution phase

synthesis of peptide nucleic acids comprising:

- (a) reacting an N-terminal protected peptide nucleic acid monomer unit with a peptide starting material to form a reaction mixture containing a peptide nucleic acid product, and
- (b) partitioning the peptide nucleic acid product from the unreacted peptide starting material, unreacted N-terminal protected peptide nucleic acid monomer unit, side-products and reagents based on the presence of

the

N-terminal protecting group.

Also claimed is the product form this reaction.

USE - The method is use for sequential solution phase synthesis of oligonucleotides and peptides

ADVANTAGE - The method lends itself to automation and is ideally suited for large scale manufacture of peptides and oligonucleotides with high efficiency. $\mathsf{Dwg.0/9}$

L12 ANSWER 8 OF 14 WPIDS COPYRIGHT 2000 DERWENT INFORMATION LTD

AN 1994-225288 [27] WPIDS

CR 1991-353169 [48]

DNC C1994-103343

TI New amino acid derivs. - useful as platelet aggregation inhibitors and in treatment of cancer.

DC B03

IN KLEIN, S I; MOLINO, B F

PA (RHON) RHONE POULENC RORER PHARM INC

CYC :

PI US 5328900 A 19940712 (199427)* 9p

ADT US 5328900 A CIP of US 1990-505286 19900405, Cont of US 1991-724675 19910702, US 1992-961216 19921015

FDT US 5328900 A CIP of US 5064814

PRAI US 1991-724675 19910702; US 1990-505286 19900405; US 1992-961216 19921015

AN 1994-225288 [27] WPIDS

CR 1991-353169 [48]

AB US 5328900 A UPAB: 19980722

Searched by John Dantzman 703-308-4488

Amino acid derivs. of formula (I), and their salts, are new X = H, amidino, COR, NR,R2,CN, NHC(=NH)R1, C(=NR,)NHR2, or a op. of formula (i) or (ii): Y = OR, NR,R2, a D-or L-amino acid (or its corresp. carboximide),

NR,CR3R4R5, NHCH(R5)-V,-CHR3R4, or a gp. of formula (iii): R1,R2 = H, alkyl, aryl, arylalkyl or alkyl; R3 = H, CO2H, CO2R1, CONH2, CONR,R2 or CONR6R7; R4,R5 = H, alkyl cycloalkyl, cycloalkylmethyl, TOR1, TSR1, TNR1R2, TNHC(=NH)NH)NH2, TC(=NH)NH2, TCO2R1, TCONR1R2, phenyl (substd. by X2), TCHPh2 (opt. ring substd. by X2), or T-Ar; Ar = a gp. of formula (iv)-(V1): etc. T = (CH2)p; P = 0-8; R6 + R7 = (CH2)4, (CH2)5, (CH2)6, CH2CH2OCH2CH2, CH2CH2NR,CH2 or a gp. of formula (x): X2 = H,Cl,Br,F,OR1, NO2,NR1R2, NHCOR1, SR1, 1-5C alkyl, phenyl, CO2R1, C(=NH)NH2,

NHC (=NH) NH2,

CONR6R7, CF3 or NHSO2R1; V, = C(O)NR1, (CH2)n, CH=CH, CH2NH, CH2O, CH2S or

C(o)CH2; m = o, 1 or 2; n = o, 1, 2 or 3.

The D- or L-amino acid is Asp, Arg, Ala, Asn, Cys, Gly, Glu, Gln, His, Ile, Leu, Lys, Met, Orn, Phe, Pro, Ser, Thr, Trp, Tyr or Val; R1, R2 = H or phenyl; R3 = H or CO2H; R4, R5 = H, alkyl or cycloalkyl; R6 + R7 = (CH2)4; m = 1; n = o; p = 1.

36 Cpds. (I) are specifically claimed, e.g., pyrrolidine-3-carboxyl-azetidine-2-carboxyl-aspartyl-valine and N-amidino-piperidine-4-carboxyl-piperidine-2-carboxyl-aspartyl- isoleucine.

(I) are **prepd**. by standard solid phase or **soln**.

phase peptide synthesis techniques.

USE - (I) are platelet aggregation inhibitors and may be used to treat or prevent thrombosis associated with certain disease states, such as myocardial infarction, stroke, peripheral arterial disease and disseminated intravascular coagulation. (I) may also be useful for treatment of certain cancerous diseases. Admin. is oral or parenteral. Dosage is $0.02-5 \, \text{mg/kg}$ day.

In an example, L-aspartyl-beta-t-butyl ester-L-valine-P-alkoxy

benzyl

resin ester was shaken with (S)-N-fmoc-azetidine-2-carboxylic acid (0.217g), EDC(0.128g), HOBT (0.091g) and NEt3 (0.1 ml), in DMF (10 ml), for 3 hrs. at room temp. The mixt. was filtered, washed, and the resin deriv. deprotected conventionally to give N-(S)-

azetidin-2-yl-carbonyl-L-

aspartyl-t-butyl ester-L-valine-p-alkoxybenzyl resin ester. This cpd. was shaken with N-60C-piperidine-4-carboxylic acid (0.205g), EDC (0.171g), hoist (0.091g) and NEt3 (0.1 ml), in DMF (10 ml), for 2 hrs. at room temp.

The mixt. was filtered, washed with CH2Cl2 and the prod. cleaned from the resin. Work up gave N-(2(S)-1-(piperidin-4-ylcarbonyl)) azetidin-2-ylcarbonyl)-L- aspartyl-L-valine as the trifluoroacetate salt, m.pt.

86-88

deg.C. In tests (as described in blood, 66, 946-952 (1985)), this cpd. inhibited fibrinogen mediated platelet aggregation with an ICso of 29.6 pM. Dwg.0/0

L12 ANSWER 9 OF 14 WPIDS COPYRIGHT 2000 DERWENT INFORMATION LTD

AN 1992-217015 [26] WPIDS

DNC C1992-098275

TI Prodn. of growth hormone releasing peptide - by soln.phase synthesis via new recrystallisable intermediates. DC B04 B05

Searched by John Dantzman 703-308-4488

```
STEVENSON, D
IN
     (SMIK) SMITHKLINE BEECHAM CORP
PA
CYC
PΙ
                   A1 19920611 (199226) * EN
     WO 9209620
        RW: AT BE CH DE DK ES FR GB GR IT LU NL SE
         W: AU CA JP KR US
     AU 9191664
                  A 19920625 (199239)
     PT 99654
                   A 19921030 (199247)
     ZA 9109440
                   A 19921230 (199306)
                                              23p
     EP 564587.
                  A1 19931013 (199341) EN
         R: AT BE CH DE DK ES FR GB GR IT LI LU NL SE
     JP 06503578
                   W 19940421 (199421)
     EP 564587
                   A4 19940824 (199533)
ADT WO 9209620 A1 WO 1991-US8863 19911125; AU 9191664 A AU 1991-91664
     19911125, WO 1991-US8863 19911125; PT 99654 A PT 1991-99654 19911129; ZA
     9109440 A ZA 1991-9440 19911129; EP 564587 A1 WO 1991-US8863 19911125, EP
     1992-903706 19911125; JP 06503578 W WO 1991-US8863 19911125, JP
     1992-503322 19911125; EP 564587 A4 EP 1992-903706
FDT AU 9191664 A Based on WO 9209620; EP 564587 Al Based on WO 9209620; JP
     06503578 W Based on WO 9209620
PRAI US 1990-621094
                      19901130
AN
     1992-217015 [26]
                        WPIDS
          9209620 A UPAB: 19931006
AB
     (A) solid recrystallisable peptide derivs. of formula (I)-(VI) are new:
          Z-L-Lys(Boc)-NH2 (I)
          Z-D-Phe-L-Lys(Boc)-NH2 (II)
          Z-L-Trp-D-Phe-L-Lys(Boc)-NH2 (III)
          Z-L-Ala-L-Trp-D-Phe-L-Lys(Boc)-NH2 (IV)
          Z-D-Trp-L-Ala-L-Trp-D-Phe-L-Lys (Boc) -NH2 (V)
          Boc-L-His (Boc) -D-Trp-L-Ala-L-Trp-D-Phe-L-Lys (Boc) -NH2 (VI)
          where Boc = t-butoxycarbonyl and Z = benzyloxycarbonyl.
          (B) Prodn. of the hexapeptide amide of formula (VII):
          L-His-D-Trp-L-Ala-La-Trp-D-Phe-L-Lys-NH2 (VIII)
          is effected by; (a) coupling (I) with Z-D-Phe to form (II); (b)
     removing Z and coupling with Z-L-Trp-NH2 to form (III); (c) removing Z
and
     coupling with Z-L-Ala to form (IV); (d) removing Z and coupling with
     Z-D-Trp to form (V); (e) removing Z and coupling with Boc-L-His(Boc) to
     form (VI); and (f) removing the Boc gps.
          USE/ADVANTAGE - (VII) has pituitary growth hormone releasing
activity
     and is useful for treating growth hormone deficiency. The process is
     capable of producing high-purity (VII) since each intermediate can be
     purified by recrystn. Decompsn. of Trp residues is minimised since only
     one acid treatment, in step (f) is required.
     0/0
ABEQ EP
           564587 A UPAB: 19931130
     (A) solid re-crystallisable peptide derivs. of formula (I)-(VI) are new:
          Z-L-Lys(Boc)-NH2 (I)
          Z-D-Phe-L-Lys(Boc)-NH2 (II)
          Z-L-Trp-D-Phe-L-Lys(Boc)-NH2 (III)
          Z-L-Ala-L-Trp-D-Phe-L-Lys(Boc)-NH2 (IV)
          Z-D-Trp-L-Ala-L-Trp-D-Phe-L-Lys (Boc) -NH2(V)
          Boc-L-His (Boc) -D-Trp-L-Ala-L-Trp-D-Phe-L-Lys (Boc) -NH2 (VI)
          where Boc = t-butoxycarbonyl and Z = benzyloxycarbonyl.
          (B) Prodn. of the hexapeptide amide of formula (VII):
          L-His-D-Trp-L-Ala-La-Trp-D-Phe-L-Lys-NH2 (VIII)
                   Searched by John Dantzman 703-308-4488
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is effected by; (a) coupling (I) with Z-D-Phe to form (II); (b)
     removing Z and coupling with Z-L-Trp-NH2 to form (III); (c) removing Z
and
     coupling with Z-L-Ala to form (IV); (d) removing Z and coupling with
     Z-D-Trp to form (V); (e) removing Z and coupling with Boc-L-His(Boc) to
     form (VI); and (f) removing the Boc gps.
          USE/ADVANTAGE - (VII) has pituitary growth hormone releasing
activity
     and is useful for treating growth hormone deficiency. The process is
     capable of producing high-purity (VII) since each intermediate can be
     purified by recrystallisation. Decomposition of Trp residues is minimised
     since only one acid treatment, in step (f) is required.
     ANSWER 10 OF 14 WPIDS COPYRIGHT 2000
                                             DERWENT INFORMATION LTD
     1989-108338 [15]
                        WPIDS
AN
DNC
    C1989-047932
ΤI
     Soln. phase synthesis of octa
     peptide with thymic humoral activity - by condensing protected
     tetra peptide then de protecting, providing high yield and easy to scale
     up.
DC
IN
     DECASTIGLI, R; FORINO, R; GALANTINO, M; DE, CASTIGLIONE R
PA
     (FARM) FARMITALIA ERBA SPA CARLO; (FARM) FARMITALIA ERBA SRL CARLO
CYC
     15
                   A 19890412 (198915) * EN
PΙ
     EP 311391
         R: AT BE CH DE ES FR GB GR IT LI LU NL SE
     AU 8823391
                   A 19890413 (198922)
                   A 19890522 (198926)
     JP 01128998
                   B1 19931229 (199401)
     EP 311391
                                       EN
                                               gę
         R: AT BE CH DE ES FR GB GR IT LI LU NL SE
     DE 3886655
                   G 19940210 (199407)
                   T3 19941216 (199505)
     ES 2061678
    EP 311391 A EP 1988-309306 19881006; JP 01128998 A JP 1988-252856
     19881006; EP 311391 B1 EP 1988-309306 19881006; DE 3886655 G DE
     1988-3886655 19881006, EP 1988-309306 19881006; ES 2061678 T3 EP
     1988-309306 19881006
FDT DE 3886655 G Based on EP 311391; ES 2061678 T3 Based on EP 311391
PRAI GB 1987-23484
                      19871007
     1989-108338 [15]
                        WPIDS
AN
           311391 A UPAB: 19930923
AΒ
     EΡ
     Prodn. of octapeptide of formula (I), and its pharmaceutically acceptable
     salts, comprises condensing protecting tetrapeptides (B) and (C);
     deprotecting the product (D), and opt. converting to salt; where X =
amino
     protecting gp.; Y and Y', opt. present, are COOH protecting gps.; K = OH
     or hydrazido; W = amino protecting gp.; Q = COOH protecting qp. or OH.
     Pref. K = OH; Y, Y' (the same) and Q are all protecting qps.
          USE/ADVANTAGE - (I) has thymic humoral activity. Compared with the
     known solid-phase synthesis (US 4621135), this soln. method provides
     easier scale-up and better yields, esp. no formation of the succinimidyl
     deriv. which is the main cyclic byproduct of the conventional method.
     0/0
ABEO EP
           311391 B UPAB: 19940217
     A process for preparing a peptide of the formula
H-Leu-Glu-Asp-Gly-Pro-Lys-
     Phe-Leu-OH (A) or a pharmaceutically acceptable salt thereof, which
     process comprises condensing a compound of the formula
                   Searched by John Dantzman 703-308-4488
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X-Leu-Glu(Y)-Asp(Y')-Gly-K (B) wherein X is an amino protecting group, Y

and Y' each independently represents a carboxy protecting group and K is а hydroxy or hydrazido group, with a compound of formula H-Pro-Lys(W)-Phe-Leu-Q (C) wherein W is an amino protecting group and Q represents a carboxy protecting group or a hydroxy group, with the proviso that Q must be a carboxy protecting group when K is a hydroxy group; deprotecting the resultant compound of the formula X-Leu-Glu(Y)-Asp(Y')-Gly-Pro-Lys(W)-Phe-Leu-Q (D) wherein X,Y,Y',W and Q are as defined above; and, if desired, converting the resulting peptide of formula (A) into a pharmaceutically acceptable salt thereof. Dwg.0/0 ANSWER 11 OF 14 WPIDS COPYRIGHT 2000 DERWENT INFORMATION LTD L12 1989-025699 [04] WPIDS AN DNC C1989-011415 New N-substd. guanidinium tetra phenyl borate salts - useful in synthesis TIof peptide(s), esp. contg. arginine. DC IN CALLENS, R; COLLIN, A (SOLV) SOLVAY & CIE PΑ CYC 19 ΡI EP 300518 A 19890125 (198904) * FR R: AT BE CH DE ES FR GB GR IT LI LU NL SE AU 8817790 A 19881222 (198907) 19881223 (198907) FR 2616784 Α JP 01016792 Α 19890120 (198909) PT 87748 Α 19890531 (198925) US 4923966 Α 19900508 (199023) 7p B1 19920902 (199236) EP 300518 FR R: AT BE CH DE ES FR GB GR IT LI LU NL SE DE 3874251 G 19921008 (199242) A 19930131 (199311) IL 86722 A 19931116 (199347) US 5262567 q₀ T3 19940101 (199405) ES 2043784 C 19940816 (199435) CA 1331496 B2 19971224 (199805) JP 2693493 g8 EP 300518 A EP 1988-201153 19880607; FR 2616784 A FR 1987-8695 19870619; ADT JP 01016792 A JP 1988-152079 19880620; US 4923966 A US 1988-207876 19880617; EP 300518 B1 EP 1988-201153 19880607; DE 3874251 G DE 1988-3874251 19880607, EP 1988-201153 19880607; IL 86722 A IL 1988-86722 19880613; US 5262567 A Div ex US 1988-207876 19880617, Cont of US 1990-486612 19900228, US 1992-854751 19920320; ES 2043784 T3 EP 1988-201153 19880607; CA 1331496 C CA 1988-569081 19880609; JP 2693493 B2 JP 1988-152079 19880620 DE 3874251 G Based on EP 300518; US 5262567 A Div ex US 4923966; ES 2043784 T3 Based on EP 300518; JP 2693493 B2 Previous Publ. JP 01016792 PRAI FR 1987-8695 19870619 1989-025699 [04] AN WPTDS 300518 A UPAB: 19930923 AΒ ΕP New guanidinium tetraphenylborate cpds. of formula (I) are new, where R = organic gp. contg. at least one amino gp. Specifically, R = -X-CH(NHA)-CO-Y; X, A and Y are each linear, branched or cyclic aliphatic gps. (opt. substd. and/or unsatd.), aromatic or aliphatic gps., or heterocyclic gps.; A can also be H and Y also OH or halo. Pref. X = (CH2)3; A = H, opt. substd. amino acid; benzyloxycarbonyl Searched by John Dantzman 703-308-4488

or tert. butoxycarbonyl; Y = OH or opt. substd. amino acid. In prepn., Ph4B(-)-salt and a cpd. contg. a guanidinium cpd. are reacted at 20-1:1, esp. 1:1, mole ratio, pref. in DMF at -60 to 100 deg.C. Pref. Ph4B(-)-salts are derived from N-contg. bases, e.g. Et3N; N-(m)ethylmorpholine; N-(m)ethylpiperidine; dicyclohexylamine or imidazole.

USE/ADVANTAGE - (I) are intermediates esp. in synthesis of peptides; partic. formation of (I) is used to solubilise Arg or peptides contg. free, but protonated, Arg residues. At the end of synthesis, the Ph4B(-) ion is easily displaced, e.g. by treating with water so as to release the guanidinium function and to reform the original Ph4B-salt which can be recovered for reuse.

ABEQ DE 3874251 G UPAB: 19930923

New guanidinium tetraphenylborate cpds. of formula (I) are new, where R = organic gp. contg. at least one amino gp. Specifically, R = -X-CH(NHA)-CO-Y; X, A and Y are each linear, branched or cyclic aliphatic gps. (opt. substd. and/or unsatd.), aromatic or aliphatic gps., or heterocyclic gps.; A can also be H and Y also OH or halo.

Pref. X = (CH2)3; A = H, opt. substd. amino acid; benzyloxycarbonyl or tert. butoxycarbonyl; Y = OH or opt. substd. amino acid. In prepn., Ph4B(-)-salt and a cpd. contg. a guanidinium cpd. are reacted at 20-1:1, esp. 1:1, mole ratio, pref. in DMF at -60 to 100 deg.C. Pref. Ph4B(-)-salts are derived from N-contg. bases, e.g. Et3N; N-(m)ethylmorpholine; N-(m)ethylpiperidine; dicyclohexylamine or imidazole.

USE/ADVANTAGE - (I) are intermediates esp. in synthesis of peptides; partic. formation of (I) is used to solubilise Arg or peptides contg. free, but protonated, Arg residues. At the end of synthesis, the Ph4B(-) ion is easily displaced, e.g. by treating with water so as to release the guanidinium function and to reform the original Ph4B-salt which can be recovered for reuse.

ABEQ US 4923966 A UPAB: 19930923

Use of guanidine-related cpds. comprising a tetraphenyl-borate ion in soln. phase peptide synthesis is

disclosed, the guanidine-related cpd. being of formula (I) where R is organic radical comprising at least one amine gp.

The cpds. are prepd. from a halogenated deriv. of carbamic acid and from substd. thiourea.

USE - Used as catalysts, plant protection agents and pharmaceutical dyes.

ABEQ US 5262567 A UPAB: 19940111

A new cpd. including a guanidine gp. and an unsubstd. tetraphenylborate ion is of formula (I), where A is H and Y is OH.

USE - (I) is soluble in organic solvents and is used in the soln. phase synthesis of peptides

contg. arginine and their protection and activation. Uses include catalysis, plant protection agents and pharmaceutical dyes. Dwg.0/0

- L12 ANSWER 12 OF 14 WPIDS COPYRIGHT 2000 DERWENT INFORMATION LTD
- AN 1989-008909 [02] WPIDS
- DNC C1989-004123
- TI New guanidinium tetra phenyl borate cpds. used as intermediates for peptide synthesis.
- DC B03 B05 C01 E12
- IN CALLENS, R; COLLIN, A

Searched by John Dantzman 703-308-4488

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(SOLV) SOLVAY & CIE
PA
CYC 19
    EP 297641
                   A 19890104 (198902)* FR
PΙ
        R: AT BE CH DE ES FR GB GR IT LI LU NL SE
                  A 19881222 (198907)
     AU 8817791
                   Α
                     19881223 (198907)
     FR 2616785
     JP 01016791
                  A
                     19890120 (198909)
     PT 87749
                  Α
                     19890531 (198925)
     US 4954616
                  A 19900904 (199038)
                                               6p
     EP 297641
                  B 19920122 (199204)
        R: AT BE CH DE ES FR GB GR IT LI LU NL SE
                   G 19920305 (199211)
     DE 3867927
     IL 86723
                   A 19930114 (199305)
    ES 2038740
                   T3 19930801 (199337)
    CA 1331497
                   C 19940816 (199435)
                                        FR
     JP 2693492
                   B2 19971224 (199805)
                                               7p
ADT
    EP 297641 A EP 1988-201152 19880607; FR 2616785 A FR 1987-8696 19870619;
     JP 01016791 A JP 1988-152078 19880620; US 4954616 A US 1988-207877
     19880617; IL 86723 A IL 1988-86723 19880613; ES 2038740 T3 EP 1988-201152
     19880607; CA 1331497 C CA 1988-569082 19880609; JP 2693492 B2 JP
     1988-152078 19880620
FDT ES 2038740 T3 Based on EP 297641; JP 2693492 B2 Previous Publ. JP
01016791
                      19870619
PRAI FR 1987-8696
     1989-008909 [02]
AN
                        WPIDS
           297641 A UPAB: 19930923
AB
     Guanidino tetraphenyl borates of formula (I) are new. R = an organic
     radical containing at least one amine function; R1-R5 = inorganic or
     organic groups. Specifically claimed is (I) where R = -(CH2)3-CH(NH2)-
     COOH; R2 = R4 = CF3; and R1 = R3 = R5 = H.
          In the prepn. a tetraphenyl borate, esp. one derived from an alkali
     or alkaline earth metal hydroxide is reacted with a cpd. contg. a
     guanidinic gp. The reaction may be effected in an organic solvent such as
     dimethyl formamide, chloroform, dichloromethane, or carbon
tetrachloride.
          USE - As intermediates for peptide synthesis.
     0/0
ABEQ EP
           297641 B UPAB: 19930923
     Guanidine-related cpds. comprising a tetraphenylborate ion, characterised
     in that they correspond to the general formula (I) in which R denotes an
     organic radical of general formula (II) in which X denotes a linear,
     branched or cyclic, substituted or unsubstituted, satd. or unsatd.
     aliphatic radical, contg. up to 25 carbon atoms, A denotes a hydrogen
     atom, an aliphatic or aromatic radical contg. heteroatoms or otherwise,
     such as protective gps. or activating gps., one or more amino acids
bonded
     by peptide bonds, in which certain functional qps. are substituted or
     unsubstituted by protective gps. or activating gps.; Y denotes a hydroxyl
     qp., a halogen atom, an aliphatic or aromatic radical contg. or not
contq.
     heteroatoms, such as protective gps. or activating gps., an amino gp., an
     amino acid or a peptide in which certain functional gps. are substituted
     or unsubstituted by protective gps. or activating gps. and by amine gps.
     of general formula NR6R7 in which R6 and R7 independently of each other
     denote a hydrogen atom or an alkyl gp. numbering from 1 to 3 carbon
atoms:
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and R1, R2, R3, R4 and R5 independently of each other denote an organic Searched by John Dantzman 703-308-4488

gp. chosen from alkyl, alkoxyalkenyl or alkenyl radicals numbering from 1

to 10 carbon atoms and contg. or not contg. heteroatoms or a hydrogen atom, at least one of the radicals R1, R2, R3, R4 and R5 being other than a hydrogen atom. ABEQ US 4954616 A UPAB: 19930923 Use of quanidine-related cpds. including tetraphenylborate ion of formula (I) in soln. phase peptide synthesis , is new. In (I) R is organic gp. contg. at least one amine gp. and opt. carboxylic gp., both opt. substd.; R1-R5 are each inorganic or organic gps. ADVANTAGE - In peptide synthesis, dissolves prod., providing activation and protection. Readily recycled. ANSWER 13 OF 14 WPIDS COPYRIGHT 2000 DERWENT INFORMATION LTD AN 1986-061717 [09] WPIDS DNC C1986-026317 TI Biologically active polypeptide recovery - by covalently binding to peptide which is capable of complexing with metal ion chelated to resin. DC B04 D16 IN PIDGEON, C; SMITH, M C (ELIL) LILLY & CO ELI PA CYC 16 A 19860211 (198609) * US 4569794 PI7p A 19860611 (198624) EP 184355 ΕN R: BE CH DE FR GB IT LI NL SE A 19860612 (198631) AU 8550240 A 19860705 (198633) JP 61148197 HU 39462 19860929 (198645) T A 19860606 (198708) DK 8505352 A 19890418 (198920) CA 1252948 IL 77104 A 19900319 (199021) EP 184355 B 19920108 (199203) R: BE CH DE FR GB IT LI NL SE DE 3585147 G 19920220 (199209) B 19930728 (199336) HU 208025 JP 07088400 B2 19950927 (199543) 10p B 19970811 (199739) DK 171917 ADT US 4569794 A US 1984-678602 19841205; EP 184355 A EP 1985-308471 19851121; JP 61148197 A JP 1985-263595 19851122; HU 208025 B HU 1985-4464 19851122; JP 07088400 B2 JP 1985-263595 19851122; DK 171917 B DK 1985-5352 19851120 HU 208025 B Previous Publ. HU 39462; JP 07088400 B2 Based on JP 61148197; DK 171917 B Previous Publ. DK 8505352 PRAI US 1984-678602 19841205 1986-061717 [09] ΑN WPIDS AB 4569794 A UPAB: 19930922 (1) A biologically active polypeptide or protein (I) covalently linked either directly or indirectly to an immobilised metal ion chelating peptide is new. (2) Recovery of (I) from the complex by selective elution with a low pH buffer. The metal ion is immobilised on a chelating resin. Protein or polypeptide recovered may be natural or synthetic and if synthetic can be prepd. by classical solution phase synthesis, solid phase synthesis or by recombinant DNA methodology, pref. the latter. They include insulin A chain, insulin B chain, proinsulin, growth hormone, glucagon, somatostatin, growth hormone releasing factor. USE - The complex is an intermediate in the recovery of the Searched by John Dantzman 703-308-4488

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biologically active polypeptide or protein (I).
     0/0
ABEQ EP
           184355 B UPAB: 19930922
     A compound comprising a biologically active polypeptide or protein
     covalently linked to a peptide that is able to chelate an immobilized
     divalent metal ion and that has two to five amino acid residues, at least
     one of which is selected from the group consisting of histidine and
     cysteine.
     ANSWER 14 OF 14 WPIDS COPYRIGHT 2000
                                             DERWENT INFORMATION LTD
     1979-00326B [01]
ΑN
                        WPIDS
TI
     Radioactively labelled calcitonin hentriaconta-peptide analogue - for
     calcitonin radioimmunoassay determination.
DC
     B04 K08 S03 S05
IN
     KUMAHARA, Y; OKADA, Y; SAKAKIBARA, S
PA
     (DARA) DAIICHI RADIOISOTOPE LAB LTD
CYC
PΙ
     DE 2826844
                   A 19781221 (197901) *
                   A 19790124 (197909)
     JP 54009293
                  A 19790223 (197913)
     FR 2395254
                  A 19810505 (198128)
     CA 1100486
                   A 19810707 (198130)
     US 4277393
                   B 19810723 (198131)
     DE 2826844
                  B 19831109 (198348)
     JP 58050213
PRAI JP 1977-73130
                      19770620
AN
     1979-00326B [01]
                        WPIDS
AΒ
          2826844 A UPAB: 19930901
     New radioiodine-labelled peptide is the hentriacontapeptide of formula
(I)
     labelled with radioactive iodine:
          Also new is the use of radioiodinated (I) as tracer in the
     radioimmunoassay of calcitonin.
          Radioiodinated (I) is more stable and purer than radioiodinated
     natural calcitonin, due to the absence of disulphide bonds, but behaves
in
     practically the same way as human calcitonin in antigen-antibody
     reactions.
          Examples describe the prepn. of the hentriacontapeptide (I)
     by solution-phase peptide synthesis
     , the radioiodination of (I) with Na125 I in the presence of chloramine
Τ,
     the prodn. of calcitonin antibody using (I) as antigen, and the use of
     radioiodinated (I) and the antibody in the radio-immunoassay
determination
     of serum calcitonin.
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